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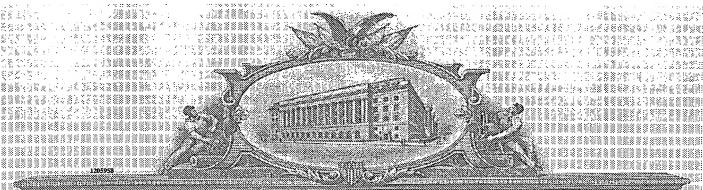
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Mail Stop Provisional Patent Application

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PROVISIONAL APPLICATION COVER SHEET

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LAST NAME							
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GROTZINGER	Joachim		Altwittenbek, Germany				
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U.S. PATENT APPLICATION

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Invention: NEW IL-11 MUTEINS

Document Name:4411-2 Title Cover Sheet Document #: 760984

Author_Id: BJS

TITLE OF THE INVENTION: new IL-11 muteins

BACKGROUND OF THE INVENTION:

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Human interleukin-11 (hIL-11) is a multi-potential cytokine that is involved in numerous biological activities such as hematopoiesis, osteoclastogenesis, neurogenesis and female fertility. It also displays anti-inflammatory properties. hIL-11 is clinically used to treat chemotherapy-induced thrombocytopenia.

Interleukin-11 was cloned from the primate stromal cell line PU-34 and was initially considered as a baematopoietic cytokine. It was later found that it also has effects on non-haematopoietic systems and that it acts on many different cell types and tissues. Numerous experiments on animal models and clinical trials with patients suffering from acute and chronic inflammatory diseases, including rheumatoid arthritis, inflammatory bowel disease, inflammatory liver disease, mucositis and psoriasis have revealed that IL-11 is an anti-inflammatory and mucosal protective agent, which, by inhibiting nuclear translocation of nuclear factor-kB (NF-kB), can reduce the production of pro-inflammatory cytokines secreted by macrophages such as TNF-α, IL-1β, IL-6 and IL-12. Its radio-protective and septic shock-protective activities have also been demonstrated in other experiments. The clinical application of hIL-11 has been approved by the FDA for the treatment of chemotherapy-induced thrombocytopenia due to the ability of this cytokine to stimulate megakaryocytopoiesis and thrombopoiesis. Another potential therapeutic application of IL-11 in the treatment of mild hemophilia A or von Willebrand disease was recently evidenced by the fact that IL-11 is able to increase von Willebrand factor and factor VIII production in a von Willebrand disease mouse model as well as in healthy mice.

Because of its broad spectrum of action, improved agonists as well as IL-11 antagonist would be of interest for numerous biological and clinical applications.

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Some structure studies of IL-11 molecule have been conducted to elucidate the interactions involved in IL-11 activation and signalling.

The structure study of Czupryn et al. 1995 thus describes the production of 61 mutated forms of hIL-11 from E. coli as thioredoxin fusion proteins [Czupryn et al. (1995) Alanine-scanning mutagenesis of human interleukin-11: identification of regions impurtant for biological activities. Ann. New York Acad. Sci. Jul 21; 762, 152-164]. Testing of these mutated forms in a murine T10 plasmacytoma proliferation assay led to the conclusion that mutations made several positions proximal to the hIL-11 C-terminus, such as a D186A mutation, caused substantial reduction in biological activity (the D186A mutation induced a 500-fold decrease in biological activity on the murine plasmacytoma cell line T10), and that a number of other mutations in this region affected either protein folding or stability.

Tacken et al. 1999 have built a three-dimensional model of human IL-11 [Tacken et al. (1999) Definition of receptor binding sites on human interleukin-11 by molecular modelling-guided mutagenesis. Eur. J. Biochem. 265, 645-655]. Three receptor binding sites within the IL-11 molecule have thus been defined (see site 1, site II and site III on Figure 1B of Tacken et al. 1999).

In Tacken et al. 1999 study, ten surface-exposed amino acid have thus selected within sites I, II and III as candidate for single point mutagenesis assays (only one amino acid per molecule has been mutated). The single point mutations made consisted in replacing a hydrophobic side chain by a charged group (aspartic acid), and a charged chain by an oppositely charged residue (lysine or glutamic acid) in order to introduce a substantial disturbance into the receptor binding sites. Nine of the ten single point mutants thus produced, including those four for which the single point mutation was on an amino acid belonging to site I (A84D mutant; L85D mutant; R190E mutant; L194D mutant), led to a substantially reduced affinity for the IL-11 receptor complex, and to a loss or a substantially reduced bioactivity (loss or substantial decrease in induction of α 1-antichymotrypsin synthesis in HepG2 cells, and of proliferation of Ba/F3-130-11 α cells). Only one of the mutants, namely R135E, which results from the replacement of a site II hydrophilic amino acid by a still hydrophilic but oppositely-charged amino acid, appeared to potentially constitute a hyperagonistic IL-11 mutant.

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There thus remains a need for a method to efficiently produce IL-11 agonists, and to obtain IL-11 agonists that would prove to be active in vivo.

SUMMARY OF THE INVENTION:

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The inventors have designed and produced IL-11 muteins wherein the hydrophobicity at site I has been substantially increased by replacement of at least two IL-11 site I hydrophilic amino acids by hydrophobic counterparts. The muteins have been characterized in terms of structure, affinity, specificity and bioactivity. Electrophoretic analysis, gel filtration, infrared spectroscopy and circular dichroism indicate that these new proteins are more compact than wild-type IL-11.

The IL-11 muteins of the invention bind to IL-11R α with an enhanced affinity (a three-fold enhanced affinity has been measured) and retain the ability to recruit gp130 through site II.

As an advantageous feature, they retain the ability to induce in vitro proliferation of various IL-11 dependent cells. A mutein of the invention; namely the H182V+D186A hIL-11 mutein, has further been shown to be 60 to 400 fold more active than wild-type IL-11 on the in vitro proliferation of 7TD1 murine hybridoma cells.

The muteins of the invention also advantageously retain in vivo biological activity. Their in vivo biological activity can further be much higher than wild-type IL-11. An injection of the H182V+D186A hIL-11 mutein at a 10-fold lower dose than the wild-type hIL-11 has been shown to delay the death of irradiated mice for the same duration.

The muteins of the invention are therefore useful in every biological, medical or clinical application in which wild-type IL-11 is useful, and can even show an enhanced efficiency. The muteins of the invention are more particularly useful in radioprotection (e.g. radioprotection of the small intestine during abdominal irradiation), in decreasing chemotherapy deleterious effects (e.g. during 5-fluoroUracil chemotherapy), in anti-inflammatory therapy, in resistance to septic shock, and in hematopoiesis stimulation.

BRIEF DESCRIPTION OF THE DRAWINGS:

Figure 1 is a reprint of AY207429 accession entry from the NCBI website (http://www.ncbi.nlm.nih.gov/entrez) giving the nucleotide and amino acid wild-type human IL-11 (hIL-11) sequences and characteristics (SEQ ID NO:73, and SEQ ID NO:1, respectively).

Figure 2 shows the complete wild-type human, macaque, mouse and rat IL-11 amino acid sequences (SEQ ID NO:1-4).

Figure 3 shows the wild-type human, macaque, mouse and rat IL-11 amino acid sequences deleted from the first 34 N-terminal amino acids (SEQ ID NO:5-8). H182 and D186 are underlined.

Figure 4 shows hIL-11 muteins of the invention (SEQ ID NO:9-13), which derive from the 34aa-deleted wild-type hIL-11 by replacement of the wild-type H182 and D186 by hydrophobic amino acids (shown underlined).

5 Figure 5 shows hIL-11 muteins of the invention (SEQ ID NO:14-18), which derive from wild-type hIL-11 deleted from its first 21 amino acids, by replacement of the wildtype H182 and D186 by hydrophobic amino acids (shown underlined).

Figure 6 shows hIL-11 muteins of the invention (SEQ ID NO:19-23), which derive from complete wild-type hIL-11, by replacement of the wild-type H182 and D186 by hydrophobic amino acids (shown underlined).

Figure 7 shows IL-11 muteins of the invention (SEQ ID NO:24-28), which derive from 34aa-deleted wild-type macaque IL-11, by replacement of the wild-type H182 and D186 by hydrophobic amino acids.

Figure 8 shows IL-11 mateins of the invention (SEQ ID NO:29-33), which derive from the wild-type macaque IL-11 deleted from the first 21 N-terminal amino acids, by replacement of the wild-type H182 and D186 by hydrophobic amino acids.

Figure 9 shows IL-11 muteins of the invention (SEQ ID NO:34-38), which derive from complete wild-type macaque IL-11, by replacement of the wild-type H182 and D186 by hydrophobic amino acids.

Figure 10 shows IL-11 muteins of the invention (SEQ ID NO:39-43), which derive from the wild-type mouse IL-11 deleted from the first 34 N-terminal amino acids, by replacement of H182 and D186 by hydrophobic amino acids (shown underlined).

- Figure 11 shows IL-11 muteins of the invention (SEQ ID NO:44-48), which derive from the wild-type mouse IL-11 deleted from the first 21 N-terminal amino acids, by replacement of H182 and D186 by hydrophobic amino acids (shown underlined).
- Figure 12 shows IL-11 muteins of the invention (SEQ ID NO:49-53), which derive from the complete wild-type mouse IL-11, by replacement of H182 and D186 by hydrophobic amino acids (shown underlined).
 - Figure 13 shows IL-11 muteins of the invention (SEQ ID NO:54-58), which derive from the wild-type rat IL-11 deleted from the first 34 N-terminal amino acids, by replacement of H182 and D186 by hydrophobic amino acids (shown underlined).
- Figure 14 shows IL-11 muteins of the invention (SEQ ID NO:59-63), which derive from the wild-type rat IL-11 deleted from the first 21 N-terminal amino acids, by replacement of H182 and D186 by hydrophobic amino acids (shown underlined).
 - Figure 15 shows IL-11 muteins of the invention (SEQ ID NO:64-68), which derive from the complete wild-type rat IL-11, by replacement of H182 and D186 by hydrophobic amino acids (shown underlined).
 - Figure 16A shows the joined CDS sequence (SEQ ID NO:69) for human complete wild-type IL-11, as defined in AY207429 NCBI nucleotide entry, and the joined CDS sequence (SEQ ID NO:70) for the blL-11 muteins of the invention which derive from the 34aa-deleted hlL-11.
- Figure 16B shows the joined CDS sequence (SEQ ID NO:71) for the hIL-11 muteins of the invention which derive from the 21aa-deleted hIL-11, and the joined CDS sequence (SEQ ID NO:72) for the hIL-11 muteins of the invention which derive from the complete hIL-11.
 - Figure 17 shows the AY207429 NCBI entry nucleotide sequence mutated in accordance with the present invention (codons n₁n₂n₃ and n₄n₅n₆ which replace the wild-type cac and gac are shown underlined).
 - Figure 18 shows the mRNA sequence (SEQ ID NO:75) of a mutein of the invention, which derives from hIL-11(codons $n_1n_2n_3$ and $n_4n_5n_6$ are shown underlined).
- Figure 19 shows the gene sequence (SEQ ID NO:76) of a IL-11 mutein of the invention which derive from hlL-11 (codons n₁n₂n₃ and n₄n₅n₆ are shown underlined).

Figure 20 shows the % of mouse survival par days of exposure to irradiation at 15 Gy (upper curve = mice treated with 3.2 micrograms of recombinant but non-mutated IL-11; lower curve = non-treated control mice).

Figure 21 shows the % of mouse survival per days of exposure to irradiation at 15 Gy

5 (upper curves = mice treated with 3.2 micrograms of recombinant but non-mutated IL
11, or with 0.32 microgram of a H182V+D186A mutein of the invention ..."HVDA"-;
lower curves = mice treated with 0.32 microgram of recombinant but non-mutated IL
11, or non-treated control mice).

Figure 22 shows the parental (non-mutated) nucleotide sequence (SEQ ID NO:77) of a recombinant IL-11 (FPΔIL-11), and its parental (non mutated) amino acid sequence (SEQ ID NO:78).

Figure 23 shows the nucleotide sequence of FPAIL-11 mutated in accordance with the present invention (SEQ ID NO:79), and the mutated corresponding amino acid sequence (SEQ ID NO:80 of the invention)

15 Figure 24 shows the primers used for inverse PCR mutagenesis of FPAIL-11.

Figures 25A and 25B show a human wild-type II-11 3D-model.

In figure 25A, a 3D model of the IL-11 based on cristallographic data obtained for CNTF, as described by Tacken et al [1999] is shown. Figure 25B shows a site I view of the IL-11 model. Positively charged amino acids (Arg, Lys) are coloured in blue, negatively charged (Asp, Glu) are in red, hydrophilies in grey and hydrophobic in yellow.

Figure 26 shows the expression of FPΔIL-11 and of the H182V+D186A mutein analysed by SDS-PAGE.

BL21 E. coli were transformed with pET-22b(1) vector encoding FPAIL-11 and H/V-D/A mutein or empty vector (E). After induction (i, induced; n, not induced) of protein production, bacteria were lysed as described in Experimental. Supernatants (100 µg of total protein per lane) were then analysed by SDS-PAGE and colored by Coomassie

Figure 27 shows infrared spectra of FPΔIL-11 (top) and of H182V+D186A (bottom) in the 1800-1400 cm⁻¹ frequency range.

The absorbance is reported in mOD. The absorbance scale is given for the bottom spectrum. The upper spectrum has been offset for clarity.

Figure 28 shows the CD spectra of the FPAIL-11 (top) and of H182V+D186A (bottom).

Figures 29A and 29B show the evolution of the integrated intensity of the amide II band as a function of the time of exposure to $^{2}H_{2}O$ for FP Δ IL-11 (circles) and for H182V+D186A (crosses).

Fitting was carried out with a three exponential decay. Panel 29A: between 0 and 20 min, panel 29B: between 0 and 700 min.

Figures 30A and 30B show gel-filtration chromatography of parental FPΔIL-11 and of mutant H182V+D186A, and their bioactivity tested from fractions collected during the chromatography.

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In Figure 30A, Superdex-75 column (K16, Pharmacia Biotech) was used and calibrated with three proteins albumin (67 kDa), ovalbumin (43 kDa) and chymotrypsinogen A (25 kDa) before loading 30 μ g of each analysed unlabelled protein in the presence of 50 ng of the same ³³P-labelled one as a tracer. Fifty microlitres of each collected fraction was submitted to a radio-counting. In Figure 30B, IL-11 activity was measured using the mouse hybridoma cell line 7TD1. Cells were cultivated in flat-bottom microwell plates (2 x 10^3 of 7TD1 cells/well) in the presence of 0.2 μ l of each eluted fraction. After 7 days of culture, the number of surviving cells was determined by colorimetric assays for hexosaminidase. Each sample was tested in triplicate and presented on average with a standard deviation.

Figures 31A and 31B show the bioactivity of parental FPAIL-11 and mutant H182V+D186A, tested on 7TD1 (Figure 31A) and on B9 cells (Figure 31B).

Cells were cultivated in flat-bottom microwell plates (2 x 10³ of 7TD1 cells/well; 1 x 10⁴ of B9 cells/well) in the presence of serial dilutions of parental FPAIL-11, mutein H/V-D/A, or commercial rhIL-11 (R&D). After 7 days for 7TD1 and 3 days for B9 cells culture, the number of surviving cells was determined by colorimetric assays for hexosaminidase (7TD1 cells) and for XTT (B9 cells). Each sample was tested in triplicate and presented as average with a standard deviation.

Figure 32 shows the inhibition of 7TD1 cells proliferation stimulated by FPAIL-11 or mutant H182V+D186A, by anti-hIL-11 and anti-human gp130 neutralizing antibodies.

Cells were incubated with the indicated concentrations of anti-human IL-11 monoclonal antibodies H2 (circles) and anti-human gp130 monoclonal antibodies MAB628

(squares) and B-R3 (triangles). Data points represent the means of triplicate determinations.

Figure 33 shows the expression of parental FPΔIL-11 and of its H182V+D186A mutein analysed by SDS-PAGE and immunoblutting.

5 BL21 E. coli were transformed with empty vector (mock) or expression vector encoding parental FPΔIL-11 or mutated proteins as indicated in the figure. After induced expression of the proteins, cells were lysed by sonication and lysates (100 μg of total protein per lane) were analysed by SDS-PAGE (left) and immunoblotting (right) using a polyclonal antibody raised against IL-11 (BAF 218).

Figure 34 shows the proliferation of 7TD1 cells in response to FPΔIL-11 and its H182V+D186A mutein.

7TD1 cells were incubated in the presence of serial dilutions of E.coli lysate containing mock, FP Δ IL-11 or muteins, which were previously adjusted to 2 μ g/ml. After 7 days of culture, the number of cells was determined by a colorimetric assay for hexosaminidase.

DETAILED DESCRIPTION:

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IL-11 signalling at present time known to be dependent on the formation of a ligand/receptor complex which comprises IL-11, IL-11R α subunit (responsible for the specificity of interaction) and gp130 receptor β -subunit (responsible for signal transduction). The interaction between IL-11 and its receptor α -subunit occurs at its recently assigned site I (Tacken *et al.* 1999, cited *supra*, and incorporated herein by reference).

Activity of IL-11 requires binding to α receptor subunit (IL-11Rα) that provides ligand specificity in a functional multimeric signal-transduction complex with gp130, the common receptor subunit for the cytokine family including IL-6, vIL-6, CNTF (Ciliary Neurotrophic Factor), LIF (Leukaemia Inhibitory factor), OSM (Oncostatin M), CI-1 (Cardiotrophin) and NNT-1/BSF-3 (Novel neurotrophin-1/B cell-stimulating factor-3).
 It is believed that IL-11 first interacts with IL-11Rα with a low affinity (K_d = 10 nM) and that the IL-11/IL-11Rα complex interacts subsequently with gp130 to form a high affinity (K_d= 300-800 pM) and signal-transducing complex.

Three sites, responsible for the interaction with the receptor subunits have been assigned for IL-11 [Grotzinger, J., Kurapkat, G., Wollmer, A., Kalai, M. and Rose-John, S. (1997) The family of the IL-6-type cytokines: specificity and promiscuity of the receptor complexes. Proteins 27, 96-109; Tacken, I., Dahmen, II., Boistcau, O., Minvielle, S., Jacques, Y., Grotzinger, J., Kuster, A., Horsten, U., Blanc, C., Montero-Julian, F. A., Ileinrich, P. C. and Muller-Newen, G. (1999) Definition of receptor binding sites on human interleukin-11 by molecular modelling-guided mutagenesis. Eur. J. Biochem. 265, 645-6551.

Site I, constituted of amino acids at the end of the AB-loop and the C-terminal part of the D-helix, is implicated in the interaction with the IL11Rα subunit. Site II, constituted of amino acids from the A and C helices and site III, constituted of the N-terminal part of the D-helix and residues from the beginning of the AB-loop, are responsible for gp130 (β-subunit) recruitment (Figure 25).

The inventors found that IL-11 muteins can be produced that have an increased affinity for IL-Ra, that have retained affinity for gp130, and that have retained or improved IL-11 biological activity.

The inventors demonstrate that such agonistic IL-11 muteins can be obtained by substantially increasing the hydrophobicity of IL-11 site I, which thereby makes the structure of the IL-11 molecule more compact. Increasing hydrophobicity of IL-11 site I can be achieved by replacement of IL-11 site I hydrophilic amino acids by hydrophobic counterparts. It further appeared to the inventors that at least two of such hydrophilic amino acids should be replaced by hydrophobic counterparts.

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The inventors notably demonstrate that site I of human wild-type IL-11 comprises two hydrophilic amino acids (His182 and Asp186), and that substituting both of them by hydrophobic counterparts (e.g. substituting His182 and Asp186 by Valine and Alanine, respectively) leads to a hIL-1 mutein with increased affinity for IL-11Rα, increased specificity and increased in vitro and in vivo bioactivity.

The fact that the muteins of the invention have such excellent properties and effects is all the more surprising and unexpected since opposite properties and effects are obtained when only one of these two hydrophilic amino acids is substituted.

- Indeed, Czupryn et al. 1995 (cited supra) describes that substituting D186 by A (without substituting H186) results in a human IL-11 mutein which has, with respect to the wild-type human II-11, a highly decreased biological activity: a 500 fold decrease in biological activity has been measured on murine plasmacytoma cell line T10.
 - This prior art hence disclosed the H/V mutation as an highly undesired candidate to obtain a mutein with increased biological activity.
- But, the inventors now demonstrate that one cannot directly rely on cell line in vitro results to reliably assume that a given mutation is a good or a bad candidate to obtain a mutein with increased efficiency: the same D186A mutation, but on a FPAIL-11 protein (Flag tag + deletion of N-terminal proline-rich region, see the examples below), has been assayed by the inventor on another cell line (cell line 7TD1, see the examples below), and appears to induce an increase in biological activity for this cell line.
 - It can also be noted that a mutein of the invention for which both His182 and Asp186 are mutated (by Val and Ala, respectively) is surprisingly and unexpectedly a lot more biologically active than human wild-type IL-11: a 60 to 400 fold increase in *in vitro* cell proliferation has been measured on 7TD1 murine hybridoma cells.
- Furthermore, as a very advantageous and in fact highly essential feature, the muteins of the invention induce an increase of biological activity in vivo in an animal, which is a mammal (a 10-fold increase in radioprotection in vivo efficiency has been measured on irradiated mice with H/V-D/A, see the examples below).
- 25 The present invention thus provides with a method to produce an IL-11 agonist, which comprises producing an IL-11 mutein wherein site I hydrophobicity has been increased by replacement of at least two non-hydrophobic amino acids which are part of the wild-type IL-11 epitope for II-11Ra by hydrophobic ones.
- 30 As said two non-hydrophobic amino acids are part of the wild-type IL-11 epitope for IL-11Ra, they belong to what is known as IL-11 site I (= end of AB-loop and C-terminal part of the D-helix).

Said at least two non-hydrophobic amino acids most preferably are surface-exposed.

The mutein molecule is thereby rendered more compact.

It has retained the ability to bind to IL-Ra through its mutated site I, and has also retained the ability to bind to the other components of the IL-11 signal transducing complex, and notably to gp130 through site II and site III of the matein.

It has also retained the ability to induce in vitro proliferation of IL-11 dependent cells, such as 7TD1 murine hydridoma cells available from the ICLC (Interlab Cell Line Collection of the Istituto Nazionale per la Ricerca sul Cancro; L.go R. Benzi, 10; 16132 Genova, Italy; see http://www.icle.it/Lista.html and http://www.biotech.ist.unige.it; ICLC Catalogue accession number = HYL96001).

It has also retained in vivo bioactivity, such as a.g. the ability to protect against radiation.

As the muteins of the invention have at least retained IL-11 affinity and bioactivity, they can be referred to as IL-11 agonist or byperagonist.

To determine whether a given IL-11 amino acid is or not part of the epitope for IL-11Rα, and whether it is or not surface-exposed, the person of ordinary skill in the art can proceed in line with what is described in Tacken et al. 1999 (cited supra). It may e.g. comprise the use of a three-dimensional structure representation of said IL-11 to locate said given amino acid so that it can be determined whether it belongs or not to site I (= epitope for IL-11Rα) and whether it is or not surface-exposed (see Figures 1A and 1B of said Tacken et al. 1999, as well as the section within this publication which is entitled "Generation of a molecular model of interleukin-11 and selection of amino acid residues for site-directed mutagenesis", the content of which is herein incorporated by reference).

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Nucleotide and amino acid sequences of wild-type IL-11Ra are available from standard sequence databanks known to the person of ordinary skill in the art. Human IL-11Ra

sequences are thus available from the NCBI website at http://www.ncbi.nlm.nih.gov/entrez under the nucleotide accession number Z38102, the content of which is herein incorporated by reference. IL-11R\alpha sequences from animal yet non-human origin, such as from mouse and rat, are also available from the NCBI website at http://www.ncbi.nlm.nih.gov/entrez (for mouse and rat IL-11R\alpha sequences, see under the respective nucleotide accession numbers X98519 and AF347936, the contents of which are herein incorporated by reference).

As a matrix to assay for binding to IL-11Ra, soluble IL-11Ra, e.g. the human IL10 11Ra-IL-2 fusion protein which is described in Blanc et al. 2000, can be used (Blanc et al. (2000) Monoclonal antibodies against the human interleukin-11 receptor alpha-chain (IL-11Ralpha) and their use in studies of human mononuclear cells. J. Immunol. Methods 241, 43-59, the content of which is herein incorporated by reference). Murine IL11-11Ralpha is available from R&D Sytems (http://www.RnDSystems.com).

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Wild-type IL-11 nucleotide and amino acid sequences are available from standard sequence databanks known to the person of ordinary skill in the art: human wild-type IL-11 sequence is described on the NCBI website at http://www.ncbi.nlnu.nih.gov/entrez under the nucleotide accession number AY207429, the content of which is herein incorporated by reference. Human wild-type nucleotide and amino acid sequences are also shown printed from said site on the enclosed Figure 1 (SEQ ID NO:73 = AY207429 IL-11 nucleotide sequence; SEQ ID NO: 1 = human wild-type IL-11 amino acid sequence). A wild-type hlL-11 cDNA sequence is also available from Accession Number NM57765 from the above-mentioned NCBI website.

Wild-type IL-11 sequences from animal yet non-human sources are also available from the above-mentioned NCBI website, such as e.g. mouse and rat IL-11 (nucleotide accession number NM 008350 and NM 133519, respectively).

The replacement of said at least two amino acids can be achieved by any standard procedure known to the person of ordinary skilled in the art. It may e.g. involve mutation by inverse PCR amplification as described in Stemmer W. P. and Morris S. K. 1992 [Stemmer and Morris (1992) Enzymatic inverse PCR: a restriction site

independent, single-fragment method for high-efficiency, site-directed mutagenesis. Biotechniques 13, 214-220], of which content is herein incorporated by reference.

The choice of appropriate primers is made by making use of the common knowledge in the field applied to the design of oligonucleotides which have such a sequence that they can have a function of primer for a given IL-11 template sequence, while having the ability to introduce at least two point mutations in the amplicon with respect to the template sequence (see Stemmer and Morris 1992, cited supra).

An illustrative procedure of such an IL-11 mutagenesis is described in example 1 below.

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The production of the mutein can be achieved by any conventional procedure known to the person of ordinary skill in the art of producing proteins in general, and of producing wild-type IL-11 in particular. It may e.g. comprise the production of a plasmid comprising a sequence coding for the mutein (for the construction of a plasmid, see Wang, X. M., Wilkin, J. M., Boisteau, O., Harmegnies, D., Blanc, C., Vandenbussche, P., Montero-Julian, F. A., Jacques, Y. and Content, J. (2002) Engineering and use of 32P-labelled human recombinant interleukin-11 for receptor binding studies. Eur. J. Biochem. 269, 61-68, the content of which is herein incorporated by reference), and transforming a host cell such as E. coli with this plasmid so that the the mutein is expressed by the transformed cells, from which it can be recovered and isolated. An illustrative procedure of mutein production is described in example 1 below.

Non-hydrophobic amino acids (e.g. hydrophilic amino acids) have a side chain that is electrically charged, or that is an uncharged yet polar chain. They notably comprise:

- Cystein (symbol = C or Cys),
 - Tyrosine (symbol = Y or Tyr),
 - Histidine (symbol = H or His),
 - Lysine (symbol = K or Lys),
 - Arginine (symbol = R or Arg),
 - Glutamine (symbol = Q or Gln),
 - Asparagine (symbol = N or Asn),
 - Glutamic acid (symbol = E or Glu),

- Aspartic acid (symbol = D or Asp),
- Serine (symbol = S or Ser),
- Threonine (symbol = T or Thr).
- Hydrophobic amino acids have a side chain that is non-polar and uncharged. They notably comprise:
 - Valine (symbol = V or Val),
 - Alanine (symbol = A or Ala),
 - Proline (symbol = P or Pro),
 - Leucine (symbol = L or Leu),
 - Isoleucine (symbol = I or Ile),
 - Methionine (symbol = M or Met),
 - Tryptophan (symbol = W or Trp),
 - Phenylalanine (symbol = F or Phe).

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Human IL-11 site I is composed of a hydrophobic cluster which comprises a limited number of hydrophilic amino acids: these site I hydrophilic amino acids notably comprise H in position 182 and D in position 186 (see SEQ ID NO:1 on Figure 1 and on Figure 2).

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In accordance with the present invention, histidine (H) in position 182 and aspartic acid (D) in position 186 are most preferred as wild-type hIL-11 mutation targets to be replaced by hydrophobic amino acids.

Similarity in terms of sterical hindrance, structure and/or size may in choosing those hydrophobic amino acids which are more appropriate to replace said H and D targets.

The most preferred hydrophobic amino acids for replacing IL-11 site I hydrophilic amino acids comprise valine (V) and alanine (A).

The mutein obtained by replacement of H182 by V and of D186 by A has proven to be 30 an IL-11 hyperagonist : compared to wild-type hIL-11, it has a three-fold increased affinity for IL-11Ra, while still retaining the ability to recruit gp130; it is 60 to 400 fold more active on the proliferation of the murine hybridoma cell line 7TD1, and the mutein reaches an *in vivo* in vivo radioprotection iso-effect at a ten-fold lower dose than the wild-type IL-11 (ten fold less mutein than wild-type IL-11 is needed to achieve the same *in vivo* radioprotection effect); see examples 1 and 2 below.

In macaque, mouse and rat wild-type IL-11, those hydrophilic amino acids which at present are known to belong to site I are also H182 and D186.

The N-terminal of wild-type IL-11 protein begins with a signal peptide of 21 amino acids, directly followed by a proline-rich region of 13 amino acids. These first 34 N-terminal amino acids are not necessary to IL-11 biological activity: they can therefore be deleted. Figure 3 shows the wild-type human, macaque, mouse and rat IL-11 sequences respectively deleted from their first 34 N-terminal amino acids (SEQ ID NO: 5-8, respectively).

- The present invention thus provides IL-11 muteins, the sequence of which comprises a sequence which derives from wild-type IL-11 deleted from their first 34 N-terminal amino acids, by replacement of the hydrophilic amino acids in positions 182 and 186 (positions calculated by reference to the complete wild-type sequence) by X₁ and X₂ respectively, X₁ and X₂ being chosen from the group comprising:
 - Valine (symbol = V or Val),

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- Alanine (symbol = A or Ala),
- Proline (symbol = P or Pro),
- Leucine (symbol = L or Leu),
- Isoleucine (symbol = I or Ile),
- Phenylalanine (symbol = F or Phe),
 - Methionine (symbol = M or Met), and
 - Tryptophan (symbol = W or Trp).

The present invention thus relates to IL-11 muteins, the sequence of which comprises a sequence chosen from the group comprising SEQ ID NO:9, SEQ ID NO:24, SEQ ID NO:39, SEQ ID NO:54 shown on Figures 4, 7, 10, 13, respectively (IL-11 muteins which derives from 34aa-deleted wild-type IL-11 from human, macaque, mouse and rat origin, respectively).

The present invention also encompasses those equivalent IL-11 muteins which comprise a sequence of at least 80%, preferably at least 90% identity with the above-mentioned SEQ ID NO:9, SEQ ID NO:24, SEQ ID NO:39, or SEQ ID NO:54, provided that X₁ and X₂ are as above-defined (i.e. hydrophobic amino acids), and that the resulting protein has retained the ability of wild-type IL-11 to induce proliferation of an IL-11 dependent cell line, such as e.g. the 7TD1 murine hybridoma cell line.

Illustrative and useful muteins of the invention comprise those for which X1 and X2 are V or A.

The present invention therefore more particularly encompasses those IL-11 muteins which comprise a sequence corresponding to a wild-type IL-11 deleted from those N-terminal amino acids which are not necessary to its biological activity, wherein the amino acids in positions 182 and 186 have been replaced by V and A, A and V, V and V, or A and A, respectively.

The present invention thus relates to those IL-11 muteins which comprise a sequence of SEQ ID NO:9, SEQ ID NO:24, SEQ ID NO:39, or SEQ ID NO:54, wherein X₁=V and X₂=A, i.e. to those IL-11 muteins which comprise a sequence of SEQ ID NO:10 (deriving from human IL-11), of SEQ ID NO:25 (deriving from macaque IL-11), of SEQ ID NO:40 (deriving from mouse IL-11), or of SEQ ID NO:55 (deriving from rat IL-11). These SEQ ID are shown on Figures 4, 7, 10 and 13, respectively.

The present invention also relates to those IL-11 muteins which comprise a sequence of SEO ID NO:9, SEQ ID NO:24, SEQ ID NO:39, or SEQ ID NO:54, wherein X_1 =A and X_2 =V, *i.e.* to those IL-11 muteins which comprise a sequence of SEQ ID NO:11 (deriving from human IL-11), of SEQ ID NO:26 (deriving from macaque IL-11), of SEQ ID NO:41 (deriving from mouse IL-11), or of SEQ ID NO:56 (deriving from rat IL-11). These SEQ ID are shown on Figures 4, 7, 10 and 13, respectively.

30 The present invention also relates to those IL-11 muteins which comprise a sequence of SEQ ID NO:9, SEQ ID NO:24, SEQ ID NO:39, or SEQ ID NO:54, wherein X₁=V and X₂=V, i.e. to those IL-11 muteins which comprise a sequence of SEQ ID NO:12

(deriving from human IL-11), of SEQ ID NO:27 (deriving from macaque IL-11), of SEQ ID NO:42 (deriving from mouse IL-11), or of SEQ ID NO:57 (deriving from rat IL-11). These SEQ ID are shown on Figures 4, 7, 10 and 13, respectively.

5 The present invention also relates to those IL-11 muteins which comprise a sequence of SEQ ID NO:9, SEQ ID NO:24, SEQ ID NO:39, or SEQ ID NO:54, wherein X₁=A and X₂=A, i.e. to those IL-11 muteins which comprise a sequence of SEQ ID NO:13 (deriving from human IL-11), of SEQ ID NO:28 (deriving from macaque IL-11), of SEQ ID NO:43 (deriving from mouse IL-11), or of SEQ ID NO:58 (deriving from rat IL-11). These SEQ ID are shown on Figures 4, 7, 10 and 13, respectively.

Illustrative and useful IL-11 muteins of the invention thus comprises those IL-11 muteins which derive from a wild-type IL-11 by deletion of the signal peptide (first 21 N-terminal amino acids), and by replacement of the amino acids in positions 182 and 186 (positions calculated by reference to the complete wild-type IL-11) by the hydrophobic X₁ and X₂ amino acids as above-defined.

The present invention thus encompasses those IL-11 muteins, the sequence of which comprises or consists in a sequence of SEQ ID NO:14, SEQ ID NO:29, SEQ ID NO:44 or SEQ ID NO:59, wherein X₁ and X₂ are as above-defined. These SEQ ID are shown on Figures 5, 8, 11, and 14, respectively.

The sequence of SEQ 1D NO:14 corresponds to the human wild-type IL-11 wherein the amino acids in positions 182 and 186 have been replaced by X_1 and X_2 , and wherein the first 21 N-terminal amino acids have been deleted (see Figure 5).

The sequence of SEQ ID NO:29 corresponds to the macaque wild-type IL-11 wherein the amino acids in positions 182 and 186 have been replaced by X_1 and X_2 , and wherein the first 21 N-terminal amino acids have been deleted (see Figure 8).

The sequence of SEQ ID NO:44 corresponds to the mouse wild-type IL-11 wherein the amino acids in positions 182 and 186 have been replaced by X_1 and X_2 , and wherein the first 21 N-terminal amino acids have been deleted (see Figure 11).

The sequence of SEQ ID NO:59 corresponds to the rat wild-type IL-11 wherein the amino acids in positions 182 and 186 have been replaced by X₁ and X₂, and wherein the first 21 N-terminal amino acids have been deleted (see Figure 14).

When $X_1=V$ and $X_2=A$ in SEQ ID NO:14, SEQ ID NO:29, SEQ ID NO:44, SEQ ID NO:59, respectively, the muteins of the invention comprise or consist in a sequence of SEQ ID NO:15, SEQ ID NO:30, SEQ ID NO:45, SEQ ID NO:60, respectively (shown on Figures 5, 8, 11, 14, respectively).

When X₁=A and X₂=V in SEQ ID NO:14, SEQ ID NO:29, SEQ ID NO:44, SEQ ID NO:59, respectively, the muteins of the invention comprise or consist in a sequence of SEQ ID NO:16, SEQ ID NO:31, SEQ ID NO:46, SEQ ID NO:61, respectively (shown on Figures 5, 8, 11, 14, respectively).

When X₁=V and X₂=V in SEQ ID NO:14, SEQ ID NO:29, SEQ ID NO:44, SEQ ID NO:59, respectively, the muteins of the invention comprise or consist in a sequence of SEQ ID NO:17, SEQ ID NO:32, SEQ ID NO:47, SEQ ID NO:62, respectively (shown on Figures 5, 8, 11, 14, respectively).

When X₁=A and X₂=A in SEQ ID NO:14, SEQ ID NO:29, SEQ ID NO:44, SEQ ID NO:59, respectively, the muteins of the invention comprise or consist in a sequence of SEQ ID NO:18, SEQ ID NO:33, SEQ ID NO:48, SEQ ID NO:63, respectively (shown on Figures 5, 8, 11, 14, respectively).

The muteins of the invention which comprise or consist in a sequence of SEQ ID NO:15-18, SEQ ID NO:30-33, SEQ ID NO:45-48, SEQ ID NO:60-63, respectively are preferred muteins of the invention. Those muteins of the invention which comprise or consist in a sequence of SEQ ID NO:15, SEQ ID NO:30, SEQ ID NO:45, SEQ ID NO:60, respectively are most preferred.

Illustrative and useful IL-11 muteins of the invention are also those muteins of the invention that derive from complete wild-type IL-11 by replacement of the amino acids in positions 182 and 186 by the hydrophobic X_1 and X_2 amino acids as above defined. Such illustrative and useful IL-11 muteins thus comprise those that comprise or consist in a sequence of SEQ 1D NO:19, SEQ ID NO:34, SEQ ID NO:49, or SEQ ID NO:64, wherein X_1 and X_2 are as above defined.

The sequence of SEQ ID NO:19 corresponds to human complete wild-type IL-11 wherein H182 and D186 have both been replaced by X₁ and X₂ as above defined. It is shown on Figure 6.

The sequence of SEQ ID NO:34 corresponds to macaque complete wild-type IL-11 wherein H182 and D186 have both been replaced by X_1 and X_2 as above defined. It is shown on Figure 9.

The sequence of SEQ ID NO:49 corresponds to mouse complete wild-type IL-11 wherein H182 and D186 have both been replaced by X₁ and X₂ as above defined. It is shown on Figure 12.

The sequence of SEQ ID NO:64 corresponds to rat complete wild-type IL-11 wherein H182 and D186 have both been replaced by X_1 and X_2 as above defined. It is shown on Figure 15.

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When $X_1=V$ and $X_2=A$ in SEQ ID NO:19, SEQ ID NO:34, SEQ ID NO:49, SEQ ID NO:64, respectively, the muteins of the invention comprise or consist in a sequence of SEQ ID NO:20, SEQ ID NO:35, SEQ ID NO:50, SEQ ID NO:65, respectively (shown on Figures 6, 9, 12, 15, respectively).

When X₁=A and X₂=V in SEQ ID NO:19, SEQ ID NO:34, SEQ ID NO:49, SEQ ID NO:64, respectively, the muteins of the invention comprise or consist in a sequence of SEQ ID NO:21, SEQ ID NO:36, SEQ ID NO:51, SEQ ID NO:66, respectively (shown on Figures 6, 9, 12, 15, respectively).

When X₁=V and X₂=V in SEQ ID NO:19, SEQ ID NO:34, SEQ ID NO:49, SEQ ID NO:64, respectively, the muteins of the invention comprise or consist in a sequence of SEQ ID NO:22, SEQ ID NO:37, SEQ ID NO:52, SEQ ID NO:67, respectively (shown on Figures 6, 9, 12, 15, respectively).

When X₁=A and X₂=A in SEQ ID NO:19, SEQ ID NO:34, SEQ ID NO:49, SEQ ID NO:64, respectively, the muteins of the invention comprise or consist in a sequence of SEQ ID NO:23, SEQ ID NO:38, SEQ ID NO:53, SEQ ID NO:68, respectively (shown on Figures 6, 9, 12, 15, respectively).

The muteins of the invention which comprise or consist in a sequence of SEQ ID NO:20-23, SEQ ID NO:35-38, SEQ ID NO:50-53, SEQ ID NO:65-68, respectively are preferred muteins of the invention. Those muteins of the invention which comprise or consist in a sequence of SEQ ID NO:20, SEQ ID NO:35, SEQ ID NO:50, SEQ ID NO:65, respectively are most preferred.

The present invention also encompasses any nucleic acid, such as DNA or RNA, coding for a mutein of the invention.

It notably encompasses any nucleic acid, such as DNA, which comprises the joined 5 CDS (coding sequences) of a wild-type IL-11 appropriately mutated in accordance with the present invention.

The joined CDS sequence of human wild-type IL-11 are shown as SEQ ID NO:69 on Figure 16A (codon CAC coding for H182, and codon GAC coding for D186 are underlined).

Appropriate mutations in accordance with the present invention comprise replacing said cac and gac wild-type codons by codon n₁n₂n₃ and n₄n₅n₆ respectively, wherein both n₁n₂n₃ and n₄n₅n₆ code for hydrophobic amino acids, *i.e.* the above-defined X1 and X2 amino acids.

Accordingly, $n_1n_2n_3$ and $n_4n_5n_6$ are both chosen from the group comprising the nucleotide codons which code for Valine (symbol = V or Val), Alanine (symbol = A or Ala), Proline (symbol = P or Pro), Lencine (symbol = L or Leu), Isoleucine (symbol = I or Ile), Phenylalanine (symbol = F or Phe), Methionine (symbol = M or Met), and Tryptophan (symbol = W or Trp). It follows that, having taken into account the degeneracy of the genetic code, $n_1n_2n_3$ and $n_4n_5n_6$ are both chosen from the group comprising the following codons:

- GCT, GCC, GCA, GCG,
- GTT, GTC, GTA, GTG,
- TTA, TTG, CTT, CTC, CTA, CTG,
- ATT, ATC, ATA,
- 25 TTT, TTC,
 - ATG,
 - CCT, CCC, CCA, CCG,
 - TGG.

The present invention thus notably encompasses any nucleic acid (e.g. DNA) which comprises the sequence of SEQ ID NO:72 shown on Figure 16B, wherein n₁n₂n₃ and n₄n₅n₅ are as above-defined.

When, in accordance with the present invention, the wild-type IL-11 has been deleted from its first 21 N-terminal amino acid (see SEQ ID NO:14-18 on Figure 5), or from its first 34 N-terminal amino acid (see SEQ ID NO:9-13 on Figure 4), the corresponding joined CDS sequence is deleted from the corresponding codons.

5 The present invention thus encompasses any nucleic acid (e.g. DNA) which comprises the sequence of SEQ ID NO:71 or of SEQ ID NO:70 shown on Figure 16D and 16A, respectively, wherein n₁n₂n₃ and n₄n₅n₆ are as above-defined.

The present invention thus more particularly encompasses any nucleic acid (e.g. DNA) which comprises or consists in the sequence of SEQ ID NO:76 or of SEQ ID NO:74, which are shown on Figures 19 and 17, respectively, and wherein the codons n₁n₂n₃ and n₄n₅n₆ are as above-defined.

The sequence of SEQ ID NO:76 corresponds to the human IL-11 wild-type gene (as defined in AY207429 NCBI accession sequence by position 1582 to position 7566), appropriately mutated in accordance with the present invention, i.e. wherein the wild-type codons cae and gae coding for H182 and D186 have been replaced by n₁n₂n₃ and n₄n₅n₆ as above-defined.

The sequence of SEQ ID NO:74 corresponds to the nucleotide sequence of SEQ ID NO:73 (AY207429 NCBI sequence), appropriately mutated in accordance with the present invention, *i.e.* wherein the wild-type codons cac and gac coding for H182 and D186 have been replaced by n₁n₂n₃ and n₄n₅n₆ as above-defined.

Following the same mutational scheme, similarly-mutated sequences can be obtained from wild-type IL-11 non-human DNA, such as macaque, mouse and rat wild-type IL-11 DNA, and such similarly-mutated sequences are encompassed by the present invention.

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The present invention also encompasses any nucleic acid comprising or consisting a RNA sequence deriving from a wild-type IL-11 RNA sequence appropriately mutated in accordance with the present invention, *i.e.* wherein the wild-type codons CAC and GAC coding for H182 and D186 have been replaced by $n_1n_2n_3$ and $n_4n_5n_6$ as above-defined.

The present invention thus particularly relates to the sequence of SEQ ID NO:75, shown on Figure 18. The sequence of SEQ ID NO:75 corresponds to the mRNA sequence of human wild-type IL-11 (as defined in AY207429 NCBI accession sequence by joined sequence 1582-1651, 3014-3186, 3386-3472, 3584-3745, 5778-7566), appropriately mutated in accordance with the present invention (n₁n₂n₃ and n₄n₅n₆ are underlined). The present invention also relates to any mutated RNA sequence which is similarly obtained from wild-type non-human IL-11 RNA, such as macaque, mouse, rat IL-11.

According to a further aspect of the present invention, the application also relates to any transfection vector, such as a.g. a plasmid, which comprises a nucleic acid of the present invention, i.e. a nucleic acid coding for an IL-11 mutein of the invention.

Illustrative and useful transfection vectors of the invention comprise those that comprise as an insertion sequence a sequence comprising or consisting of a sequence which derives from a sequence coding for a wild-type II-11 by replacement of the codons coding for the hydrophilic amino acids in positions 182 and 186 by the codons n₁n₂n₃ and n₄n₅n₆ as above-defined, and possibly by deletion of codons that are not necessary to an biological activity of the IL-11 type, such as e.g. by deletion of the codons which in the complete wild-type IL-11 code for the N-terminal signal peptide and/or the those N-terminal codons corresponding to proline-rich regions.

Illustrative and useful transfection vectors of the invention thus can comprise a sequence comprising or consisting of a sequence which derives from a sequence coding for a wild-type IL-11 by replacement of the codons coding for the hydrophilic amino acids in positions 182 and 186 by the codons n₁n₂n₃ and n₄n₅n₆ as above-defined, and by deletion of those codons of the complete wild-type IL-11 that code for the first 21 N-terminal amino acids or for the first 31 or 34 N-terminal amino acids.

A short nucleotide sequence coding for a Flag tag, such as Asp-Tyr-Lys-Asp-Asp-Asp-Asp-Asp-Lys, followed by another short nucleotide sequence coding for a consensus sequence that can be recognized and phosphorylated by a kinase (such as Arg-Arg-Ala-Ser-Val-Ala that can be recognized and phosphorylated on the serine residue by the bovine heart kinase) can be added at one end of the IL-11 mutein encoding nucleic acid, e.g. at the 5' part of it, in lieu et place of the codons which in the complete wild-type IL-11 code for the first 31 N-terminal amino acids.

Such a transfection vector is described in example 1 below. An illustrative and useful insertion sequence for such a vector is shown on Figure 23 under SEQ 1D NO:79 (wild-type human IL-11 which has been mutated in accordance with the present invention by the above-defined n₁n₂n₃ and n₄n₅n₆ codons, which has been deleted from the codons coding for the first 31 N-terminal amino acids, and to which codons coding for a Flag tag and for consensus sequence recognized by the heart bovine kinase have been added: in Figure 23, the Flag tag is boxed, and the consensus sequence for kinase is underlined).

According to a further aspect of the invention, the present application relates to any cell comprising a nucleic acid of the invention, and/or which has been transfected by a transfection vector of the invention, and/or which express a mutein of the invention.

Such cells may e.g. be used to produce and isolate IL-11 muteins of the invention. Any cell that is available to the person of ordinary skill in the art as appropriate host cell may used to be transformed by a transfection vector of the invention, so that the transformed cell thereby produced can express a mutein of the invention. Appropriate standard host cells e.g. comprise E. coli cells, such as e.g. the E. coli BL21(DE3) strain (available from Novagen).

The invention thus encompasses a method to produce an IL-11 mutein of the invention which comprises culturing a cell of the invention in a suitable culture medium (e.g. for a E. coli transformed cell, in Luria-Bertani medium), and isolating said IL-11 mutein from said cell.

The present invention thus encompasses any cell transformed with a nucleic acid sequence of the invention in operative association with an expression control sequence capable of directing replication and expression of said nucleic acid sequence.

As indicated above and illustrated below, the IL-11 muteins of the invention have at least retained an ability to bind to IL-11Ra and gp130, and to induces an *in vitro* and *in vivo* activity of the type induced by wild-type IL-11. They thus are all useful in every application in which wild-type IL-11 is considered as useful.

Exemplary biological or medical applications of IL-11 are described in US 6,126,933; WO 00/74707; US 5,460,810; US 6,540,993; US 5,215,895; WO 00/53214; the contents of which are herein incorporated by reference.

The IL-11 muteins of the invention for which H182 and D186 have been replaced by Val and Ala further prove to be highly more efficient than wild-type IL-11, and may thus be referred to as IL-11 hyperagonists. For example, the H182V+D186A muteins of the invention (i.e. the muteins which comprise a sequence of SEQ ID NO:10, SEQ ID NO:25, SEQ ID NO:40, or SEQ ID NO:55, or conservative variants thereof) bind to IL-11Rα with a three-fold enhanced affinity compared to wild-type IL-11, are 60 to 400 fold more active on the proliferation of 7TD1 murine hydridoma cells than wild-type IL-11, and are required at a ten fold lower dose to induce the same in vivo radioprotection wild-type IL-11 iso-effect (see examples 1 and 2 below).

The present application therefore also encompasses the IL-11 muteins of the invention as agents useful for improving resistance to radiation, such as resistance to radiation therapy for the treatment of cancer or for the preparation of patients for bone marrow transplantation.

The present application also encompasses the IL-11 muteins of the invention as agents useful for improving resistance to deleterious effects induced by chemotherapy for the treatment of cancer.

It more particularly relates to the IL-11 muteins of the invention as antithrombocytopenia agents.

The IL-11 muteins of the invention can also be useful as anti-inflammatory agents, and/or as agents to induce or stimulate hematopoiesis, neurogenesis, osteoclastogenesis, and/or female fertility.

The present invention thus relates to any drug that comprises a therapeutically effective amount of an 1L-11 mutein of the invention, or a nucleic acid of the invention, or a vector of the invention, or a cell of the invention. Such a drug may further comprise any pharmaceutically-acceptable vehicle (e.g. isotonic sodium chloride solution) that is available to the person of ordinary skill in the art of preparing drugs, as well as any stabilizer, preservative, buffer, antioxidant, or additive that the person of ordinary skill

will find appropriate. The drug may be produced in any form and conditioning that is appropriate to its intended mode of administration (parenteral, intravenous, subcutaneous, topical, etc.). The dosage regimen of a drug of the invention will be determined by the attending physician considering the condition, body weight, sex, diet, age, and other medically-relevant features of the patient to be treated. As an advantageous feature of the invention, those drugs which comprise the muteins for which H182 and D186 by Val and Ala will be usually required at lower doses than would have been wild-type IL-11.

0 A drug of the invention may be useful for stimulating and/or enhancing cells involved in the immune response and cells involved in the proper functioning of the hematopoietic system.

It may also be useful for treating inflammatory bowel diseases (e.g. Crohn's disease, ulcerative colitis, indeterminate colitis and infectious colitis), mucositis (e.g. oral mucositis, gastrointestinal mucositis, nasal mucositis, and proctitis), necrotizing enterocolitis, inflammatory skin disorders (e.g. psoriasis, atopic dermatitis, and contact hypersensitivity), aphtous ulcers, pharyngitis, esophagitis, peptic ulcers, gingivitis, periodontitis, and ocular diseases (e.g. conjunctivitis, retinis, and uveitis).

It may also be useful to prevent or treat hemorrhagic shock, and to protect the gastroinstestinal system during a hemorrhagic shock and resuscitation.

It may also be useful to prevent or treat an immune-mediated cytotoxicity, such as graft versus host disease or rejection of organ and tissue transplants, as well as non-immune-mediated necrotic injuries, such as localized tissue or cell injury caused by loss of blood supply, corrosion, burning, or the local lesion of a disease.

25 The invention more particularly relates to any anti-thrombocytopenia drug, which comprises a mutein of the invention, and a therapeutically effective amount of an active principle for chemotherapy of cancer.

The following examples are given as illustrative examples, and are in no way intended
to restrict the scope of the present invention. More particularly, while human IL-11
muteins are described as an useful and particularly relevant illustration, any

conservative variants thereof that the person of ordinary skill in the art will contemplate are encompassed by the present application.

EXAMPLE 1: production of IL-11 muteins, and characterization of their structure, affinity, specificity and cell line bioactivity.

EXPERIMENTAL

Bacterial strains, enzymes and chemicals

Escherichia coli DH5a was from Invitrogen Life Technologies. BL21 (DE3) and pET
22b(+) were from Novagen. E. coli recombinant human IL-11 was from PeproTech Inc. (London, UK) and R & D Systems (Wiesbaden-Nordenstadt, Germany). Primers for mutagenesis were from Genset. MAB628 and anti-hIL-11 biotinylated polyclonal antibody BAF218 were from R & D Systems. [\gamma^{-32}P]ATP with a specific radioactivity of ~3000 Ci/mmol was obtained from Amersham Pharmacia Biotech. Acrylamide and N,N'-methylene-bisacrylamide were from Bio-Rad. RPMI-1640, DMEM, glutamine and FCS were from Gibco-BRL. The catalytic subunit of cAMP-indepentent protein kinase from bovine heart muscle, streptavidin conjugated alkaline phosphatase, sodium dodecyl sulfate (SDS) and anti-Flag M2 monoclonal antibody were obtained from Sigma (Bornem, Belgium).

20 Mutagenesis

FPAIL-11 was mutated by inverse PCR amplification of the plasmid pET-FPAIL-11 previously described in Wang et al. 2002, using the primers shown on Figure 24, followed by a *Dpn* I digestion to eliminate the parental plasmid.

For a detailed description of the mutation by inverse PCR amplification, see Stemmer, W. P. and Morris, S.K. (1992) Enzymatic inverse PCR: a restriction site independent, single-fragment method for high-efficiency, site-directed mutagenesis. Biotechniques 13, 214-220, the content of which is herein incorporated by reference.

For the construction of plasmid pET-FPAIL-11, see Wang, X. M., Wilkin, J. M., Boisteau, O., Harmegnies, D., Blanc, C., Vandenbussche, P., Montero-Julian, F. A., Jacques, Y. and Content, J. (2002) Engineering and use of 32P-labelled human

recombinant interleukin-11 for receptor binding studies. Eur. J. Biochem. 269, 61-68, the content of which is herein incorporated by reference.

Please note, in accordance with Wang et al. 2002, the N-terminal nucleotides encoding the first 31 amino acids of human IL-11 joined CDS shown on Figure 16A under SEQ ID NO:69 have been deleted (the first 21 signal peptide amino acids + the 10 amino acids which follow and which correspond to a proline-rich region), and replaced by a sequence encoding a Flag tag (Asp-Tyr-Lys-Asp-Asp-Asp-Asp-Lys) followed by a consensus amino-acid sequence (Arg-Arg-Ala-Ser-Val-Ala) that can be recognized and phosphorylated on the serine residue by the bovine heart kinase. FPAIL-11 therefore has the following sequence (upper nucleotide line in italic = human IL-11 joined CDS sequence SEQ ID NO:69, lower nucleotide line = FPAIL-11; bottom amino acid line = FPAIL-11 protein):

ATG AAC TGT GTT TGC CGC CTG GTC CTG GTC GTG CTG AGC CTG

15

M D Y K D D D D K E G R

25

The Flag tag of FPAIL-11 is boxed; the phosphorylation site recognized by the bovine heart protein kinase catalytic subunit created in FPAIL-11 is underlined.

Hence, the nucleotide sequence of the non-mutated parental FPΔIL-11 is the sequence of SEQ ID NO:77, and its amino acid sequence is the sequence of SEQ ID NO:78, shown on Figure 22.

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And, the nucleotide sequence of FPAIL-11 mutated in accordance with the present invention is the sequence of SEQ ID NO:79, and the amino acid protein sequence of this mutated FPAIL-11 is the sequence of SEQ ID NO:80, shown on Figure 23.

Gel purified PCR fragments were ligated overnight at 16°C using T4 DNA ligase and then used to transform E. coli DH5α. The corresponding plasmids were amplified in DH5α, sequenced and then used to transform the BL21(DE3) strain of E. coli.

Production and purification of parental and mutant FPAIL-11

BL21 (DE3) cells transformed with the plasmid carrying the mutant or parental 15 FPΔIL-11 cDNA were cultured in Luria-Bertani medium containing 100 μg/ml of ampicillin. Expression of the recombinant proteins was induced by 1mM IPTG for 2h at 37°C.

E. coli were then lysed by 30 minutes incubation at 37°C in presence of 0.1% triton X-100 and 150 µg/ml of lysozyme in 50 mM Hepes, pH 7.4, followed by sonication for 5 minutes at an intensity level of 5 using a microprobe (Vibra Cell, Sonics Materials Inc. Danburg, Connecticut, USA). Lysates were centrifuged two times at 13,000 g for 25 min at 4°C and then assayed or purified as previously described [Wang et al. 2002, cited supra, and incorporated by reference]. Briefly, lysates were precipitated with 60% (NH4)2SO4 in order to concentrate the crude proteins. Salts were eliminated by dialysis against 50 mM Hepes, pH 7.4 buffer before the purification of samples by chromatography on a Mono-S HR5/5 column (Amersham Pharmacia Biotech) using a 50 mM Hepes buffer, pH 7.4, and a 0-1 M NaCl gradient.

Quantification of parental and mutant FPAIL-11 by ELISA

Two antibodies raised against human IL-11, a non-neutralising monoclonal antibody MAB618 and a biotinylated polyclonal BAF218, were used to quantify the recombinant human parental and muteins by sandwich ELISA method. 96-wells plates were coated overnight at 4°C with 100 µl of monoclonal antibody MAB618 at a concentration of 2 µg/ml. After blocking with 3% BSA, 100 µl of serial dilution of samples were added and incubated for 1 hour at 37°C. After washing with PBST buffer (PBS buffer in the presence of 0.1% of Tween 20), plates incubated for another hour at 37°C with 100 µl/well of biotinylated polyclonal antibody BAF218 at a concentration of 30 ng/ml. Before another incubation at 37°C for 1 hour with straptavidin conjugated alkaline phosphatase (1/5000), the plates were washed 3 times with TBS buffer (100 mM Tris-HCl, 150 mM NaCl, pH 7.5). Finally, the test was revealed using an ELISA Amplification System (Gibco BRL). Commercial recombinant IL-11 was used as a standard and the sensitivity was 2 pg/ml.

15 Mass Spectrometry

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The exact molecular weight of the FPAIL-11 and the mutein was determined using nano-electrospray mass spectrometry on a hybrid quadrupole Time-of-Flight Q-TOF mass spectrometer (Micromass, Whytenshawe, UK). Prior to analysis, samples were desalted using Vivaspin microconcentration devices with a cut-off of 10 kDa (Millipore, Bedford, MA). After washing twice with water, samples were dissolved in a mixture of 50 % acetonitrile and 0.1 % formic acid in water to a concentration of approximately 5 pmol/µl. Four µl of this sample were loaded in a nano-electrospray capillary (MDS Proteomics, Odense, Dk) that was then placed in the special holder delivered with the instrument. Spray was initiated by slightly breaking the needle tip and supplementing a small back-pressure of nitrogen. The capillary voltage was set at 1250 V. Spectra were accumulated for about 5 minutes, collecting data from m/Z 1000 to 2500 at 1 sec per scan. Data processing was performed using the Masslynx and MaxEnt software delivered with the instrument.

Infrared spectrometry

ATR-FTIR spectra were recorded at room temperature on a Bruker IFS55 FTIR spectrophotometer equipped with a liquid nitrogen-cooled mercury-cadmium-telluride (MCT) detector at a nominal resolution of 2 cm⁻¹ and encoded every 1 cm⁻¹. The internal reflection element (IRE) was a germanium plate (50×20×2mm) with an aperture angle of 45°, yielding 25 internal reflections. The spectrophotometer was continuously purged with air dried on a FTIR purge gas generator 75-62 Balston (Maidstone, England) at a flow rate of 10-20 Umin in the sample compartment and 5 I/min in the optic compartment. Thin films were obtained by slowly evaporating a sample under a stream of nitrogen on one side of the ATR plate [Fringeli and Gunthard (1981). Infrared membrane spectroscopy. Mol. Biol. Biochem. Biophys. 31, 270-332, the content of which is herein incorporated by reference]. The ATR plate was then sealed in a liquid sample holder. The sample on the ATR plate was rehydrated by flushing ²H₂O-satured N₂, at room temperature. 256 scans were averaged for each measurement. Secondary structure determination was based on the shape of the amide I band (1600-1700 cm⁻¹), which is sensitive to the secondary structure [Goormaghtigh et al. (1990). Secondary structure and dosage of soluble and membrane proteins by attenuated total reflection Fourier-transform infrared spectroscopy on hydrated films. Eur. J. Biochem. 193, 409-420, the content of which is herein incorporated by reference].

Hydrogen/deuterium exchange kinetics: nitrogen was saturated with ²H₂O by bubbling in a series of three vials containing ²H₂O. Before starting the deuteration, 10 spectra of the sample were recorded to test the stability of the measurements. At zero titue, the ²H₂O-saturated N₂ flux, at a flow rate of 100ml/min (controlled by a Brooks flow meter), was connected to the sample. For each kinetic time point, 24 scans were recorded and averaged at a resolution of 4 cm⁻¹. All the spectra of the kinetics were corrected for atmospheric water absorption and side chain contribution. The subtraction of atmospheric water was done automatically by a home-written software which computed the subtraction coefficient as the ratio of the atmospheric water band between 1579 and 1572 cm⁻¹ on the sample spectrum and on a reference atmospheric water spectrum [45-49]. The area of amide II, characteristic of the δ(N-II) vibration, was

obtained by integration between 1596 and 1502 cm⁻¹. For each spectrum, the area of amide II was divided by the corresponding amide I v(C=O) area. This ratio expressed in percentage was plotted versus deuteration time. The 100% value is defined by the amide II/amide I ratio obtained before deuteration. The 0% value corresponds to a zero absorption in the amide II region, observed for a full deuteration of the protein.

Circular dichroism

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CD measurements were carried out on a Jasco J-720 spectropolarimeter (Japan Spectroscopic Co., Ltd., Tokyo, Japan) equipped with a temperature control unit and calibrated according to Chen and Yang [Chen and Yang (1977). Two-point calibration of circular dichrometer with d-10-camphorsulfonic acid. Anal. Lett. 10, 1195-1207, the content of which is herein incorporated by reference]. The spectral bandwidth was 2 nm (< 250 nm) and 1 nm (> 250 nm), respectively. The measurements were carried out at a temperature of 20°C, the solvent was PBS throughout. The time constant ranged between 1 and 4 s and the cell path length between 0.1 and 10 mm.

15 Labelling of FPAIL-11 and of its mutein

FPΔIL-11 and H/V-D/A were labelled through protein phosphorylation with $[\gamma^{-12}P]$ ATP in the presence of bovine heart kinase and phosphorylation was checked by autoradiography as previously described [Wang et al. (2002), cited supra, and incorporated by reference].

20 SDS-PAGE and Western blot

SDS-PAGE was carried out as previously described [Laemmli (1970). Cleavage of structural proteins during the assembly of the head of bacteriophage T4. Nature 227, 680-685, the content of which is herein incorporated by reference]. Muteins and parental FPAIL-11 were transferred from gels to a nitrocellulose membrane and detected by incubation with biotinylated goat polyclonal antibody BAF218 (R&D), then streptavidin-conjugated alkaline phosphatase and finally revealed with the NBT/BCIP system (Sigma). Alternatively, proteins were detected using a biotinylated anti-flag antibody (M2 antibody, from Sigma).

Binding of 32P-H/V-D/A to cells

Binding of ³²P-H/V-D/A on 7TD1 cells was carried out as described for the parental ³²P-FPΔIL-11 on B13Rα1 cells by Wang *et al* [Wang *et al*. (2002), cited *supra*, and incorporated by reference]. *T*TD1 cells (5 x 10⁵) were pre-incubated in culture medium lacking growth factor for 2 h and were washed 3 times with phosphate-buffered saline, pH 7.4 (PBS). For binding studies, radiolabelled H/V-D/A was added to cells at the indicated concentration in PBS containing 0.5% bovine serum albumin. The mixture was incubated at 4 °C for the appropriate time and bound radiolabelled H/V-D/A was separated from the free radioactivity by centrifugation at 3000 g for 1 min through a 0.2 ml layer of a mixture of 40% dioctyl phthalate and 60% dibutyl phthalate (Janssen Chimica, Beerse, Belgium). After quick freezing, the tip of each tube containing the cell pellet was cut-off and radioactivity was counted in a Beckman β-counter. Non-specific binding was determined by incubating cells with radiolabelled H/V-D/A in the presence of a 200-fold molar excess of unlabelled H/V-D/A.

15 Surface plasmon resonance studies

These experiments were performed with a BiaCore 2000 optical biosensor (BiaCore, Uppsala, Sweden). A fusion protein of human IL-11R and IL-2 (IL-11R-IL-2) [Blanc et al. (2000). Monoclonal antibodies against the human interleukin-11 receptor alphachain (IL-11Ralpha) and their use in studies of human mononuclear cells. J. Immunol. Methods 241, 43-59, the content of which is herein incorporated by reference] was coupled through primary amino groups to a carboxymethyl dextran flow cell (CM5) at a low immobilisation level (about 500 RU per flow cell) compatible with kinetic binding studies. Subsequent binding of parental FPAIL-11 or mutein was carried out in Hepesbuffered saline (pH 7.4) at a flow rate of 10 µl/minute at room temperature.

25 IL-11 bioassay

IL-11 activity was measured using the 7TD1 cells. $2x10^3$ cells/well were cultured in flat-bottom 96 wells microtiter plates during 7 days in the presence of serial dilutions of the purified mutein or parental FP Δ IL-11, or *E. coli* crude lysates containing different muteins previously adjusted to the same protein concentration. The cell number in each

well was then determined by a colorimetric assay for hexosaminidase [Van Snick et al. (1986). Purification and NH2-terminal amino acid sequence of a T-cell-derived lymphokine with growth factor activity for B-cells hybridomas. Proc. Natl. Acad. Sci. U.S.A. 83, 9679-9683, the content of which is herein incorporated by reference]. Bioactivity was assayed similarly on 1 x 10⁴ B9 cells/well for about 3 days and revealed by XTT colorimetric assay. Each sample was tested in triplicate using a commercial recombinant human IL-11 (from PeproTech) as a standard.

RESULTS

Expression, purification and initial characterisation of the H/V-D/A mutein

10 FPΔIL-11 was used as the human IL-11 parental molecule to generate IL-11 muteins by mutagenesis because i) it has the same biological activity as the wild-type human recombinant IL-11 and ii) the presence of the flag-tag (F), the phosphorylation site (P) and the absence of the first ten amino acids of IL-11 (Δ) allow a strong expression, a simple purification and an easy radio-labelling of IL-11 (Wang et al. (2002), cited supra].

To evaluate the involvement of H182 and D186 residues with respect to biological activity and receptor binding, the corresponding positions were substituted by site-directed mutagenesis using an inverse PCR method [Stemmer and Morris (1992), cited supra, and incorporated by reference]. These two residues were replaced by a valine (H182/V) and an alanine (D186/A) to generate a mutein named H/V-D/A.

The expression of these parental and mutant FPAIL-11 in E. coll was analysed by SDS-PAGE (Figure 26). The parental molecule had an apparent molecular mass of about 24 kDa, a value higher than its theoretically expected one (20.050 kDa). This difference could be due to the introduction of numerous charged residues present in the 125 flag-tag and the phosphorylation site at the N-terminus of FPAIL-11 (1 Glu, 5 Asp, 2 Arg and 2 Lys). Indeed, when the two charged residues H182 and D186 of FPAIL-11 were replaced by two hydrophobic amino acids, the resulting mutein moved faster in gels than its parent, so that its apparent molecular mass (19 kDa) was close to its calculated one (19.9 kDa). This observation reinforced the hypothesis that the charged

residues could influence the molecular mobility in SDS-PAGE. However, to rule out the possibility that the reduced mobility of the H/V-D/A could be linked to a truncation of the protein, purified parental and mutant FPAIL-11 were submitted to mass spectrometric analysis. FPAIL-11 and H/V-D/A were found to have masses of 20.016 kDa and 19.934 kDa respectively, in perfect agreement with their predicted molecular masses.

Even though the increased electrophoretic mobility of the H/V-D/A mutein on SDS-PAGE is most likely due to charge modifications, we can not rule out the possibility that it would be partially due to a structural and/or conformational change of the molecule induced by mutagenesis. Such changes could render the mutein more compact than the parental molecule, therefore making it more resistant to heat denaturation and move faster in polyacrylamide gels.

Structural analysis by infrared spectrometry (IR) and circular dichrolsm (CD)

In order to further evaluate a potential conformational change induced by mutagenesis, the parent and mutant proteins purified to homogeneity were characterized by attenuated total reflection Fourier transform infrared spectrometry (ATR-FTIR). This technique has been successfully used to investigate the structure of soluble and membrane proteins [Goormaghtigh et al. (1990), cited supra]. The method is based on the analysis of the vibration bands of protein and particularly the amide I band, ν(C=O), whose absorption frequency is dependent upon the secondary structure. Figure 27 represents the ATR-FTIR deuterated spectra of those two proteins recorded at pH 7.4. Their similar spectra suggest that the replacement of two amino acids (H182 and D186) by a valine and an alanine, respectively, does not have a detectable effect upon the protein secondary structure. The main absorption peak within the amide I is located in a region associated to the α-helical structure, confirming that this structure is predominant in both IL-11 (parent and mutein).

Parental and mutant IL-11 were also submitted to CD analysis because this technique is more sensitive to α-helical structures. Figure 28 shows their CD spectra. Both spectra have the same shape but their intensity is different. Secondary structure analysis [Kalai

et al. (1997). Analysis of the human interleukin-6/human interleukin-6 receptor binding interface at the amino acid level: proposed mechanism of action. Blood 89, 1319-1333, the content of which is herein incorporated by reference] of the far UV CD spectrum of both proteins reflect the α -helical character of the proteins (parental IL-11: α -helix 44.8 %, β -sheet 14.0%, turn 15%, remainder 26.2%; mutant IL-11: α -helix 38.8%, β -sheet 17.0%, turn 15.7%, remainder 28.5%), which are typical for a four-helix bundle cytokine. The somewhat lower helical content of the IL-11 mutein compared to the parental might reflect a conformational change introduced by the mutated amino acids.

To further characterize conformational changes taking place upon mutagenesis of FPΔIL-11, deuteration kinetics of the mutein and its parental protein were measured. In a soluble protein, the rate of hydrogen/deuterium exchange is essentially related to protein structure stability (local unfolding dynamics in secondary structures govern the exchange). The hydrogen exchange rate of the proteins was followed by monitoring the amide II absorbance peak decrease [δ(N-H) maximum in the 1596-1502 cm⁻¹ region] because of its shift to the 1460 cm⁻¹ region [amide II', δ(N-D)] upon deuteration (data not shown). The variations with time of the percentages of non-exchanged residues, calculated from the ratio of amide II/amide I as described in Experimental, are shown in Figure 29. It appears that the FPΔIL-11 is undergoing a fast exchange, whereas H/V-D/A mutein is more resistant to hydrogen/deuterium exchange, suggesting that the mutein might form oligomers and/or have a more compact structure than parental FPΔIL-11.

By gel-filtration on a Superdex-75 column, parental and mutant proteins were both eluted at a similar position corresponding to a monomeric form (Figure 30), indicating that the increased hydrophobicity due to mutagenesis at site I did not lead to the formation of dimers or oligomers.

Interaction with soluble IL-11Ra

In order to find out if mutagenesis and associated conformational change of H/V-D/A have an effect on its interaction with IL-11Ra, the association and dissociation kinetic constants (k_{on}, k_{off}) describing parent IL-11 and H/V-D/A mutein binding to human

IL-11R α were determined by surface plasmon resonance biosensor analysis using dextran-immobilized purified human IL-11R α -IL-2 [Blanc *et al.* (2000), cited *supra*] fusion protein as matrix. As depicted in Table 1 below, the association (k_{on}) and dissociation (k_{off}) kinetic constants of H/V-D/A were both much higher (35 and 14 fold respectively) than those of parental FP Δ IL-11, leading to an equilibrium dissociation constant (K_{ol}) for the mutein that was 3-fold lower than for FP Δ IL-11.

Table 1: Kinetic (k_{on} association, k_{off} dissociation) and equilibrium (K_{off} dissociation) constants for the binding of FPΔIL-11 and H/V-D/A to the recombinant human IL-11R-IL-2, determined by surface plasmon resonance.

10				
	IL-11	k _{an} (M ⁻¹ s ⁻¹)	k _{off} (s ⁻¹)	K _d (nM)
	FPΔIL-11	$5.90 (\pm 0.90) \times 10^3$	9.75 (± 0.05) x 10 ⁻⁴	165 (±25)
	H/V-D/A	2.30 (± 0.74) x 10 ⁵	1.34 (± 0.46) x 10 ⁻²	58 (±1.5)

If one translates the equilibrium dissociation constants in terms of free energies of interaction ($\Delta G = -RTln(1/K_D)$), binding of FP Δ IL-11 or H/V-D/A to U-11R-II-2 is accompanied by free energy changes of 9.2 or 9.8 kcal/mol, respectively, indicating that the mutagenesis and its induced conformational change favour IL-11 interaction with the IL-11R α receptor.

Interaction with cell surface IL-11 receptors

15

B13Ra1 and 7TD1 cells were used to test H/V-D/A binding to human and murine IL-11 receptors. B13Ra1 are Ba/F3 cells stably transfected with human gp130 and hII.-11Ra [Lebeau et al. (1997). Reconstitution of two isoforms of the human interleukin-11 receptor and comparaison of their functional properties. FEBS Lett. 407, 141-147, the content of which is herein incorporated by reference]. Non-specific binding component, determined by adding a 200-fold molar excess of unlabelled H/V-D/A, was low (less than 5% of the total association). Analysis of the specific binding data by the method of Scatchard indicated the existence of a single class of binding sites (see Table 2 below).

<u>Table 2</u>: Dissociation constants and numbers of sites per cell of FP Δ IL-11 and H/V-D/A binding on B13R α 1 and 7TD1 cells

	B13Ra1		7TDI			
Ligands			Class 1 sites		Class 2 sites	
	K _e (nM)	Sites/cell	K _d (nM)	Sites/cell	K _d (nM)	Sites/cell
³² P-FPΔIL-11			1			
competed with FPAIL-11	0.44	3079	7.20	391	0.65	16
competed with H/V-D/A	0.40	2900	ND*	ND	ND	ND
³¹ P-H/V-D/A	1				 	
competed with H/V-D/A	0.71	3462	2.70	486	0.60	16
competed with FPAIL-11	0.72	3531	סא	ND	ND	ND
ND: non determined						

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We could only detect high affinity receptors on these cells probably because of an excess of gp130 expression on the surface of the transfected cells. The dissociation constant for the mutein ($K_d = 0.7 \text{ nM}$) was higher than that for its parent ($K_d = 0.4 \text{ nM}$). Binding of ³²P-H/V-D/A could be completely inhibited by an excess of FP Δ IL-11 and the reverse was also found, showing that the two molecules compete with each other for this binding.

7TD1 is a murine hybridoma cell line resulting from the fusion of the mouse myeloma cell line Sp2/0-Ag14 with spleen cells from a C57BL/6 mouse. This cell line is well known to respond to picogram amounts of IL-6 [Van Snick et al. (1986), cited supra], but has also a proliferating response to nanogram amounts of IL-11 [Wang et al. (2002), cited supra].

When 7TD1 cells were used for 32 P-labelled H/V-D/A or FP Δ IL-11 receptor binding assays, two classes of binding sites were observed (see the above Table 2): low affinity receptors with K_d in the nanomolar range likely corresponding to the binding of IL-11 or mutein to isolated IL-11R α chains, and high affinity receptors with K_d in the picomolar range likely corresponding to the association of IL-11/IL-11R α with gp130 transducing subunits. Similar numbers of either types of receptors were detected with

labelled FP Δ IL-11 and H/V-D/A, in agreement with the above observation that the two molecules compete for common receptors. In the context of low affinity binding to isolated IL-11R α chains, the affinity of H/V-D/A ($K_d = 2.7$ nM) was found to be around 3-fold higher than that determined for FP Δ IL-11 ($K_d = 7.2$ nM), in agreement with the biosensor experiments (see the above Table 1). In the context of the high affinity receptor complex however, no differences were found between H/V-D/A and FP Δ IL-11 binding ($K_d = 0.60$ nM vs 0.65 nM).

Induction of cell proliferation

To investigate to what extent the increased affinity of the mutein for the IL-11Ra could impact on its bioactivity, cell proliferation assays were conducted on different cell lines.

As shown in Figure 31A, H/V-D/A mutein, like IL-11, supports 7TD1 cell proliferation dose-dependently. However, the concentration of the mutein required to induce half-maximal proliferation (EC₅₀) was much lower (400-fold) than that required for the wild-type IL-11 (EC₅₀ = 0.03 ng/ml for H/V-D/A vs 15 ng/ml for FPAIL-11 and rldL-11). This increased activity of the mutein was consistently found in several experiments, with a H/V-D/A/FPAIL-11 activity ratio ranging from 60 to 400. Gel filtration experiments (Figures 30A and 30B) showed that parental and mutant IL-11 behaved as monomeric molecules (at about 20 kDa) with no sign of aggregation, and biological activity was fully associated with these monomers.

In sharp contrast to what was found on 7TD1 cells, the H/V-D/A mutein was about 10fold less active on B9 cells (Figure 31B), another murine hybridoma cell line, indicating
that the mode of action of the IL-11 mutein was more complex than expected.

In order to check if the H/V-D/A activity was mediated through gp130 transduction, we used an anti-IL-11 mAb (H2) that has been demonstrated to react with an epitope localized in site II of IL-11 [Blanc et al. (2000), cited supra]. By interfering with gp130 recruitment, this antibody inhibits the binding of FPAIL-11 to its receptors and consequently inhibits IL-11-dependent cell proliferation [Wang et al. (2002), cited supra]. Figure 32 shows that this neutralizing antibody is able to inhibit 7TD1 cell proliferation induced by both the parental and mutant FPAIL-11, indicating that the

epitope recognized by the antibody H2 (site II) is conserved on H/V-D/A mutein, and that H/V-D/A, like parental IL-11, requires the gp130 subunit for exerting its bio-activity. The anti-human gp130 antibodies MAB628 and B-R3 did not affect parental or mutant IL-11 proliferation of the murine 7TD1 cells, and served as controls. As far as these two antibodies have been shown to inhibit cell proliferation on human cells [Chevalier et al. (1996). Interleukin-6 family of cytokines induced activation of different functional sites expressed by gp130 transducing protein. J. Biol.Chem. 271, 14764-14772], these results also indicate that the epitopes recognized by these antibodies on human gp130 are not shared by murine gp130.

When analysing the dose-response curves depicting the inhibitory effect of H2 antibody (Figure 32), it appeared that the concentration of H2 necessary to induce half-maximal inhibition (IC₅₀) was about 10 fold lower in the case of the H/V-D/A mutein than in the case of parental IL-11. This indicates that the H/V-D/A mutations at site I induce a conformational change at site II that results in an increased affinity for the H2 antibody. Other experiments showed that H/V-D/A, like IL-11, was able to stimulate the proliferation of Ba/F3 cells co-transfected with human IL-11Rα and human gp130, whereas Ba/F3 cells only transfected with human gp130 were insensitive to either molecule. Therefore H/V-D/A, like parental IL-11, cannot activate gp130 in the absence of IL-11Rα.

20 Relative roles of H182 and D186 in the properties of H/V-D/A

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In order to investigate the relative importance of H182 and D186, these residues were mutated separately or in combination generating H/V, D/V, D/A and H/V-D/V muteins, in addition to H/V-D/A. As shown in Figure 33, SDS-PAGE and Western blot analysis indicate a good expression of all recombinant proteins. As observed before for H/V-D/A, all muteins showed systematic differences between their apparent molecular mass on SDS-PAGE and their predicted one. Muteins D/V and D/A moved faster than the mutein H/V, suggesting that the negatively charged residue (D) had more impact on the molecular mobility in gels than the positively charged one (H). The difference of mobility between D/V and D/A also indicated that the charge is not the only factor involved in the mobility change. This reinforces our previous hypothesis that beyond the charge, an SDS resistant conformational change of the molecules resulting from the

mutagenesis could also contribute to the mobility change. The two double muteins H/V-D/V and H/V-D/A had similar and higher mobilities than the single muteins, indicating cumulative effects of the two mutations.

7TD1 cells were used to measure the bioactivity of the various FPΔIL-11 muteins (Figure 34). It appeared that the D/A mutation alone resulted in a strong increase in activity, even stronger than the H/V-D/A combination. The D/V mutation also resulted in an increase in activity but to a far lower extent than the D/A mutation. In contrast, the H/V mutation always resulted in a slight reduction of bioactivity: H/V, H/V-D/V and H/V-D/A were less active than wild-type, D/V and D/A respectively.

These results suggest that D186 is a key amino acid in site I and plays an essential role in the activity of IL-11. Of note, replacement of D186 by a valine instead of an alanine resulted in a much lower increase of activity, suggesting that in addition to the hydrophobic nature, the size of the side chain at position 186 is crucial for this enhancement of activity. The H182 residue also appears to be involved in the interaction at site I but with a minor role.

DISCUSSION

The aim of this study was to create potent agonists of human IL-11 by changing amino acids located in the area (site 1) responsible for binding to the specific receptor chain (IL-11Rα). A model of IL-11 (Figure 25) was built by homology considerations based on the known receptor interaction sites of the related cytokines IL-6, CNTF, and LIF [Jacques et al. (1998). The interleukin-11/receptor complex: rational design of agonists/antagonists and immunoassyas potentially useful in human therapy. Res. Immunol. 149, 737-740, the content of which is herein incorporated by reference]. Supported by mutagenesis experiments, the model predicts that the main energy for receptor ligand binding is provided by hydrophobic interactions of a few apolar side chains shielded by a surrounding scaffold of polar or charged residues which guarantee the specificity of the interaction by the formation of hydrogen bonds and salt bridges [Kalai et al. (1997), cited supra]. Therefore, in order to enhance the interaction of IL-11 with its α-receptor subunit, we replaced two charged amino acids residues H182 and D186 located in the middle of the site I hydrophobic cluster by two hydrophobic ones.

We anticipated that increasing locally the hydrophobicity on the surface of site I could influence the quaternary structure of the molecule: a putative large hydrophobic interaction zone generated by mutagenesis might favour H/V-D/A to form oligomers. Superdex-75 chromatography has evidenced that II/V-D/A is in fact expressed as a soluble functional monomeric protein. However, IR hydrogen/deuterium exchange kinetics showed that the H/V-D/A mutein is more resistant to ¹H/²H exchange, suggesting that the mutein might have a more compact structure than parental FPΔIL-11. IR ¹H/²H kinetic studies were indeed recorded at a higher protein concentration since the proteins were concentrated in a film for that experiment. It is then conceivable that additional interactions are present in the IR experiment. Yet, such local interactions encompassing the new, more hydrophobic, domain found in the H/V-D/A mutant could not explain the large effect reported on figure 29 where almost 40% of the residues experience a slower exchange, nor can such a difference be explained in view of the minor differences in the secondary structures. On the other hand, a more compact structure is deduced from the mutant's faster mobility on SDS-PAGE, in good agreement with the slower IR 1HPH exchange and CD data.

10

Analysis of the binding characteristics of the H/V-D/A mutein confirmed that residues at the end of the D-helix are implicated in recognition for and interaction with IL-11Rα. Indeed, biosensor studies showed that the H/V-D/A mutations were associated with modifications in the parameters of binding to the isolated IL-11Rα chain. Both the association and dissociation constants were markedly increased, indicating that the nature of the molecular bonds involved in the cytokine-receptor interaction at site I were strongly modified. Despite these changes, the binding affinity of the mutein for IL-11Rα was only three-fold higher than that of parental IL-11. Equilibrium studies on cell surface receptors confirmed this three-fold increase in affinity and further showed that in the context of the high affinity IL-11Rα/gp130 complex, the mutein and wild type IL-11 displayed similar affinities.

The relative bioactivity of the H/V-D/A mutein as compared to wild type IL-11 was not correlated to the difference in affinity between the two molecules. Indeed, on the 7TD1 murine hybridoma cell line, the H/V-D/A had a considerably (up to 400-fold) increased activity, whereas on another murine hybridoma cell line (B9), its bioactivity was

reduced by about 10-fold. Such variations are in line with a previous study showing that, on another murine plasmocytoma cell line (T10), the substitution of the D186 by an alanine (D/A mutein) rendered the cytokine 500-fold less active than the wild-type [Czupryn et al. (1995), Ann. New York Acad. Sci. 762, 152-164, cited supra and herein incorporated by reference].

What makes the H/V-D/A more active on 7TD1 cells? Since 7TD1 cells are highly responding to IL-6, such a high H/V-D/A bioactivity could result from their stimulation via IL-6Ra-mediated signal transduction. As parental FPAIL-11 was found in this study to fully compete with ¹²P-labelled H/V-D/A for its high affinity binding to 7TD1 cells and since the binding of this radio-labelled protein to IL-6Ra was not detectable in a RIA assay, this hypothesis can be refuted. The induction by H/V-D/A of murine IL-6 can be also excluded since we found that H/V-D/A bioactivity on 7TD1 was not modified in the presence of an IL-6 neutralising antibody. One has therefore to hypothesize that another factor, whose expression is cell line dependent, is responsible for the enhanced activity of the H/V-D/A mutein. Such a factor could be another unknown receptor chain participating to the structure of the functional IL-11 receptors. The stoichiometry of IL-11 ligand-receptor complex is still an open question and a transducing subunit different from gp130 might participate in IL-11 mediated signal transduction. A possible candidate for this unknown subunit is the gp130-like receptor (GLM-R) that has been recently identified and found to be expressed predominantly on activated monocytes [Ghilardi et al. (2002). A novel type I cytokine receptor is expressed on monocytes, signals proliferation, and activates STAT-3 and STAT-5. J. Biol. Chem. 277, 16831-16836, the content of which is herein incorporated by reference]. This receptor is able to transduce a proliferation signal and induce activation of the transcription factors STAT-3 and STAT-5. Even though its ligand has not yet been identified, GLM-R was not found to be «per se» a receptor for IL-11.

In the frame of such a hypothesis (heterocomplex of gp130 with gp130-like receptor), one could postulate that the conformational change induced by mutagenesis could render the mutein H/V-D/A more prone than wild type IL-11 to recruit and/or activate this unknown gp130-like factor. As far as our studies on 7TD1 cells showed that H/V-D/A and wild type IL-11 displayed similar high affinity binding, the higher activity of

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H/V-D/A would be related to a higher signal transduction efficiency. Therefore, on cells that would express gp130-like in excess to gp130 (like 7TD1), the mutein would be more active, and conversely on cells (like B9 or T10) that would express gp130 in excess to gp130-like the mutein would be less active. Neutralizing antibodies inhibition experiments showed that site II of H/V-D/A remained functional, although its conformation was modified with respect to antibody I12 binding. Such a modification at site II could lead to the recruitment by H/V-D/A of a gp130-like molecule instead of gp130. Alternatively, site II of H/V-D/A would still be involved in recruitment of gp130 and site III involved in recruitment of gp130-like.

In conclusion, we have generated novel hIL-11 muteins with enhanced affinity for IL-11Ra and strongly enhanced activity on 7TD1 cells. These muteins therefore constitute agonist molecules potentially useful in pathologies in which IL-11 has been shown to be beneficial. In addition, it should be a valuable molecule in further studies aiming at precising the structure and function of the IL-11 receptors.

15

EXAMPLE 2: in vivo radioprotection.

Mice. C₅₇Bl/6 male mice, 8-12 weeks old, were purchased from Charles River Laboratories (Chatillon sur Chalaronne, France). Mice were housed at the animal core facility of INSERM U463 at Nantes (France). This facility is approved by the Préfecture of the French Department of Loire-Atlantique and is maintained in accordance with the regulations and standards of French Veterinary Services.

Radiation and II-11 treatment. Whole body irradiation was delivered with a Teratron 780 (Atomic Energy of Canada limited, Canada) operating ⁶⁰Co sources. The dose rate was 1.5 Gy/min. Human recombinant FPAII-11 and H/V-D/A proteins (synthesized by Jean Content (Institut Pasteur, Bruxelles, Belgium) was solubilized in sterile PBS containing 0.2% gelatin, and delivered intravenously by retro-orbital injection of 800 ng, 30 minutes before irradiation and 5, 60 and 120 minutes after irradiation.

Survival studies. Survival as an end point was calculated from the time of treatment until death using the product limit Kaplan-Meier. Differences in product limit Kaplan-Meier survival curves were evaluated by the Mantel log-rank test for censored data. Statistical analysis was performed by Student's t test.

Results

One of the major problems encountered by radiotherapists upon irradiation of patients' abdomen is the great radio-sensitivity of the gastrointestinal tract. Local irradiation of the abdomen leads to destruction of the intestinal villae, resulting in dehydration, septic shock and subsequently death of patient. This pathology is known as gastrointestinal syndrome (GI syndrome). It has long been established that death of the stem cells located in the intestinal crypts prevents the regeneration of epithelial cells causing necrosis of the villae (Potten CS, Merritt A, Hickman J, Hall P, Faranda A: Characterization of radiation-induced apoptosis in the small intestine and its biological implications. Int. J. Radiat. Biol. 65:71-8. 1994).

In mice, we observed that a single dose of 15 Gy induced total destruction of the intestinal mucosa and subsequent death of the animal (Paris F, Fuks Z, Kang A, Capodieci P, Juan G, Ehleiter D, Haimovitz-Friedman A, Cordon-Cardo C, Kolesnick R: Endothelial apoptosis as the primary lesion initiating intestinal radiation damage in mice. Science. 293:293-7., 2001).

We therefore evaluated the therapeutic potential of FPΔIL-11 (described in the above example 1) in lethally irradiated C57BL6/J mice exposed to γ-rays, and found that FPΔIL-11 delays the death of the animals (median death 8 days for the mice pretreated by the FPΔIL-11 and irradiated versus 5 days for the mice vehicle treated and irradiated at 15 Gy). Results are illustrated by Figure 21.

In the same experimental conditions, we have evaluated the therapeutic activity of the H/V-D/A mutated proteins. A pretreatment with a 10 time lower dose of H/V-D/A, as compared to the dose used for the FPAIL-11 (total doses $0.32 \mu g$ versus $3.2 \mu g$) delays the mortality to the same iso-effect (median death at 8 days). Results are illustrated by Figure 22. Pretreatment with FPAIL-11 at this low dose $(0.32 \mu g)$ has only a little impact in survey of the animal irradiated at 15 Gy.

These experiments show that the H/V-D/A mutein of the invention provides with a gain of function, as compared to the wild type IL-11, and improved the protection of the small intestines after exposure to radiation.

ABSTRACT

The present invention relates to new IL-11 muteins of which site I hydrophobocity has

5 been increased. These muteins act as IL-11 agonist or hyperagonist, and are notably
useful as anti-thrombocytopenia agents, and as agents improving the resistance of an
organism to the deleterious in vivo effects induced by radiation or chemotherapy during
the treatment of cancer or for the preparation of patient to transplantation.

http://www.ncbi.nlm.nih.gov/entrez

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2003
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VERSION
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  ORGANISM
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REFERENCE
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  AUTHORS
             Rieder, M.J., Carrington, D.P., da Ponte, S.H., Hastings, N.C., Ahearn, M.O., Kuldanek, S.A., Rajkumar, N., Toth, E.J., Yi, Q. and
             Nickerson, D.A.
  TITLE
             Direct Submission
  JOURNAL
             Submitted (26-DEC-2002) Genome Sciences, University of
Washington,
             1705 NE Pacific, Seattle, WA 98195, USA
COMMENT
             To cite this work please use: SeattleSNPs. NHLBI HL66682
Program
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             http://pga.gs.washington.edu/).
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3241 gaggagtotg cgggcagcca cttggagggg ttctgggctc tcaggtggca gagtgaggga 3301 ggggaagagt tggggggctg gcgtggggga tggagggagc cccgaggctg ggcaggggcc 3361 acctcacage tttttccct gccagaggga caaattccca gctgacgggg accacaacct 3421 ggattccctg cccaccctgg ccatgagtgc gggggcactg ggagctctac aggtaagggc 3481 aagggagtgg gctggggaca aggtgggagg caggcagtga agggggcggg gaggatgagg 3541 ggcactggtc gggtgttctc tgatgtcccg gctctatccc cagctcccag gtgtgctgac 3601 aaggetgega geggaeetae tgteetaeet geggeaegtg eagtggetge geegggeagg 3661 tggctcttcc ctgaagaccc tggagcccga gctgggcacc ctgcaggccc gactggaccg 3721 gctgctgcgc cggctgcagc tcctggtatg tcctggcccc aagacctgac accccagacc 3781 eccaecetg geoccaaaat cetgtggeet gagteettga ageetgagae eccagaeceg 3841 agtgcaacag coccyctety agaccetyae accetaacag cocyctetya gaccetyaea 3901 ccgtaacage eccgetetga gaccetgace etaacagtee tgetetgaga ecctgaceet 3961 gcagteccaa gatectgtgg ceetgagace etgaggeeet agaceeecaa atectgeeca 4021 gaaacttcaa attotoacco aagaccotga gactocatca tocatgacot caaagteeco 4081 agateccage cectaagace caagacecca teetgaagee caaageettg agaatteaaa 4141 tecteacete aagaettgga gaccetggee ceatgaeatt gaaaaceatg gacetggeea 4201 ggcgtggtgg ctcacgcctg taatcccagc actttgggag gccgaggcaa gtggatcacc 4261 tgaggtcggg agttcaagac cagccagacc aacatggtga aaccctgtct ctactaaaaa 4321 tacaaaatta gccaggcgtg gtggtgcatg cctgtaatcc cagctacttg ggaggctgag 4381 gcaggagaat cgcttgaacc tgggaggcgg aggttgcagt gagccgagat cgcaccatta 4501 aaaagaagga aaagaaaacc atggacctcc agaccctgag accccaggcc ccagccctga 4561 gatcctgaca tettaaagat cecaggeeet aagatacaag acettgacee aaageeagee 4621 ttgggaccct ggctgtacaa acccaagacc tccaggacct agaccccgag ccctgaggcc 4681 ctatgtctca ctcccaacat cgaaaaccct gacacctcag atcctgagcc tgcgcctgta 4741 egactecaag acceteactt ccaaagecag geccaaagec etgagaceag aagaetteaa 4801 accotggtto ttgggcotaa ctocaaagac cotggatoto aaattocaac ttotagotot 4861 gagactocag coctcaccca tgagttoctg aacttgaacc cagagacccc atetotaaga 4921 etteageett gagateeagg geetgaceet agaetegage ecacagaeet eagataetgt 4981 ctgtaaaacc ccagctctgg tggggagcag tggctcactc ctgtaatccc aaggcagggg 5041 aggccaaggc agaaggacct cttgaggcca tgagtttgag acagcctggg cagcatagca 5101 agactotgtt tottaattat tattattatt attattttt ggagacagag totogogoto 5161 tgttgcccag gctagagtgc aatggtgcca tttcggcttg ctggaacctc cgcctcctgg 5221 gctcaagcga ttctcctgcc tcagcctcct gagtagctgg gacttcaggt gcacactgcc 5281 acacceggat aattittitg tattitagta gacacagggt ticaccgtgt tgcccaggct 5341 ggtcacaaac teetgagete aggecateeg eeegeetegg eeteccaaag egetgggata 5401 acaggegtga tecegegege etggettett aattgtteta acageageea caacaacaaa 5461 aacccagete tgagatteca geeceggega etetaacagt eceaggeeeg ateceteace 5521 tagaaccgag atgccagccc tgactccaca gacttcaccc ccaaccccca cactcagctc 5581 tggaagcccg tcctgactcc agcetccatt ttcggaaccc caeagcctga agagetcccg 5641 gcctaaacac ttcaccccac gcgccacagt ccccctqtga atatgcagcc ccqattcagc 5701 tgcagctcca cagcacccct gccctgcacc cccgctgcac cccctacctg tgactcacct 5761 ctetectete eccaeagatg tecegeetgg ecctgeecca gecaeecceg gaecegeegg 5821 egececeget ggegeeeece tecteageet gggggggeat eagggeegee eacgceatee 5881 tgggggggct gcacctgaca cttgactggg ccgtgagggg actgctgctg ctgaagactc 5941 ggctgtgacc cggggcccaa agccaccacc gtccttccaa agccagatct tatttattta 6001 tttatttcag tactgggggc gaaacagcca ggtgatcccc ccgccattat ctccccctag 6061 ttagagacag tccttccgtg aggcctgggg ggcatctgtg ccttatttat acttatttat 6121 ttcaggagca ggggtgggag gcaggtggac tcctgggtcc ccgaggagga ggggactggg 6181 gtcccggatt cttgggtctc caagaagtct gtccacagac ttctgccctg gctcttcccc 6241 atctaggcct gggcaggaac atatattatt tatttaagca attacttttc atgttggggt 6301 ggggacggag gggaaaggga agcctgggtt tttgtacaaa aatgtgagaa acctttgtga 6361 gacagagaac agggaattaa atgtgtcata catatccact tgagggcgat ttgtctgaga 6421 gctggggctg gatgcttggg taactggggc agggcaggtg gaggggagac ctccattcag 6481 gtggaggtcc cgagtgggcg gggcagcgac tgggagatgg gtcggtcacc cagacagctc 6541 tgtggaggca gggtctgagc cttgcctggg gccccgcact gcatagggcc gtttgtttgt

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6661 cactgeaacc tecaceteec gggtteaagc aatteteetg ceteageete eegattaget
6721 gggatcacag gtgtgcacca ccatgcccag ctaattattt atttcttttg tatttttagt
6781 agagacaggg tttcaccatg ttggccaggc tggtttcgaa ctcctgacct caggtgatec
6841 tectgeeteg geeteecaaa gtgetgggat tacaggtgtg agecaccaca cetgacccat
6901 aggicticaa taaatattta atggaaggit ccacaagica ccctgtgatc aacagtaccc
6961 gtatgggaca aagctgcaag gtcaagatgg ttcattatgg ctgtgttcac catagcazac
7021 tggaaacaat ctagatatec aacagtgagg gttaagcaac atggtgcatc tgtggataga
7081 acgccaccca gccgcccgga gcagggactg tcattcaggg aggctaagga gagaggcttg
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7741 ggagtgggga ctotgttgca caaagtcaca cagotaggga gaggtggaag tgggattotg
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7861 acatgcacac acatgagata tggagaaaca ggttcagtaa ggatttgggt cttacccagg
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8281 aggagacggg ggttgcggtg agccgagatc acatcacaaa cagccctagg cagtgcgggg
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8461 atggagtete getetgttge etaggetgga gtgeagtgtg geaatetaag eteactgeet
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8941 gtgggtccct gttctggggg agctgcaaaa gaccctccag aagggcgagt acctgcccct
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9121 ggatttagca gtcactgtgt gggggacgat caggagggag gctcaggctg tggctgctgg
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9301 gcctggtgtt gcctgacatc ttgctgatcg gccagcccgc cgaggacagg gactgctccg
9361 gootogtgot gaccaggtgo ogcatococo aaccootogg cogococoto caccootoot
9421 getetagacg etecectete ecteteccag gatgatecce etggaceteg tecacetetg
9481 cgtccatgac ctctctgcct ggcgcctgaa gctgcgcctg gtctcgggcc gccagtacta
9541 cetggecetg gacgecettg acaacgaggt gggetteetg ttecaetget gggteegect
9601 catcaaccty cttcaggage eggeteceae etggaceeee aggaceaege geaeggeeee
9661 cctggatatg ccgctggcca aagcgcctgc ctccacctgg cacctgcagg tgggatccca
9721 gctccacaga ccagggcatg gcaggcccca ggaaccctcc ggccagatcc agaggggact
9781 cgaccaagag cccaaagtct agg
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60 120 180	60 120 180	60 120 180	60 120 180
51 ADTRQLAAQL LRRAGGSSLK IRAAHAILGG	VO:2- : 51 EDTRQLTIQL LRRAMGSSLK IRRAHAILGG	51 ADTRQLAAQM LRRAGGPSLK IRAAHAILGG	51 ADTRQLAAQM LRRAAGPSLK IRAAHAILGG
lete native human IL-11 -SEQ ID NO:1-: 1	Complete native macaque IL-11 (Macaca fascicularis) -SEQ ID NO:2-: 1 11 21 31 41 51 1 MNCVCRLVLV VLSLWPDTAV APGPPPGSPR ASPDPRAELD STVLLTRSLL EDTRQLTIQL 61 KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK 121 TLEPELGTLQ TRLDRLIRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG IRAAHAILGG	(Mus musculus) -SEQ ID NO:3-: 31 41 APGPPAGSPR VSSDPRADLD SAVLITRSLL ADTRQLAAQM SAGTLGSLQL PGVLTRIRVD LMSYLRHVQW IRRAGGPSLK QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG	ste native rat IL-11 (Rattus norvegicus) -SEQ ID NO:4-: 1 21 31 41 MNCVCRIVLV VLSIMPDRVV APGPPAGSPR VSSDPRADLD SAVILTRSIL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRIRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLIRRI QLIMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG
0:1-: 31 R VSPDPRAEL L PGVLTRLRA P QPPPDPPAP	fascicular 31 R ASPDPRAEL PSVLTRLRA	11us) -SEQ 31 R VSSDPRADLI PGVLTRLRVI	egicus) -SI 31 (VSSDPRADLI PGVLTRLRVI
1 -SEQ ID NG 21 7 APGPPPGPP1 8 SAGALGALQ1 QLLMSRLAL1	.11 (Macaca 21 7 APGPPPGSPI 1 SAGALGALQI 9 QLLMSRLALI	(Mus muscu 21 APGPPAGSPR SAGTLGSLQI QLLMSRLALE	Rattus norv 21 APGPPAGSPR SAGTLGSLQI QLLMSRLALP
ite native human IL-11 1 11 MNCVCRLVLV VLSLWPDTAV RDKFPADGDH NLDSLPTLAM TLEPELGTLQ ARLDRILRRI LHLTLDWAVR GLILLKTRI	te native macaque IL- 1	te native mouse IL-11 1	rat IL-11 (11 VLSIMPDRVV NLDSIPTIAM ARIERILRRI GLIIKTRI
Complete native human IL-11 -SEQ ID NO:1-: 1	ete native MNCVCRLVLV KDKFPADGDH TLEPELGTLQ LHLTLDWAVR	O)	Complete native rat IL-11 (Rattus norvegicus) -SEQ ID NO:4-1 21 31 41 41 1 1 21 31 41 41 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Comple 1 61 121 181	Comple 1	Comple 1 61 121 181	Comple 1 61 121 181

Native human IL-11 deleted from the 34 first aminoacids -SEQ ID NO :5-:

PRAELD STVLLTRSLL ADTRQLAAQL RDKFPADGDH NLDSLPTLAM SAGALGALQL PGVLTRLRAD LLSYLRHVQW LRRAGGSSLK TLEPELGTLQ ARLDRLLRRL QLLMSRLALP QPPPDPPAPP LAPPSSAWGG IRAAHAILGG LHLTLDWAVR GLLLKTRL

Native macaque IL-11 deleted from the 34 first aminoacids -SEQ ID NO:6-:

PRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ TRLDRLLRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG IRAAHAILGG L<u>H</u>LTL<u>D</u>WAVR GLLLKTRL

Native mouse IL-11 deleted from the 34 first aminoacids -SEQ ID NO:7-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG L<u>H</u>LTL<u>D</u>WAVR GLLLLKTRL

Native rat IL-11 deleted from the 34 first aminoacids -SEQ ID NO:8-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG L $\underline{\mathbf{H}}$ LTL $\underline{\mathbf{D}}$ WAVR GLLLLKTRL

hIL-11 mutein deriving from 34aa-deleted native human hIL-11 -SEQ ID NO :9-:

PRAELDSTVLLTRSLLADTRQLAAQLRDKFPADGDHNLDSLPTLAMSAGALGA LQLPGVLTRLRADLLSYLRHVQWLRRAGGSSLKTLEPELGTLQARLDRLLRRL QLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRAAHAILGGL \underline{X}_1 LTL \underline{X}_2 WAVRGLL LLKTRL wherein X_1 and X_2 are chosen from the group comprising :

- Alanine (A),
- Valine (V),
- Leucine (L),
- Isoleucine (I),
- Phenylalanine (F),
- Methionine (M),
- Proline (P),
- Tryptophan (W).

hIL-11 mutein deriving from 342a-deleted native human hIL-11 -SEQ ID NO :10-:

PRAELDSTVLLTRSLLADTRQLAAQLRDKFPADGDHNLDSLPTLAMSAGALGA LQLPGVLTRLRADLLSYLRHVQWLRRAGGSSLKTLEPELGTLQARLDRLLRRL QLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRAAHAILGGL<u>V</u>LTL<u>A</u>WAVRGLLL LKTRL

hIL-11 mutein deriving from 34aa-deleted native human hIL-11 -SEQ ID NO :11-:

PRAELDSTVLLTRSLLADTRQLAAQLRDKFPADGDHNLDSLPTLAMSAGALGA LQLPGVLTRLRADLLSYLRHVQWLRRAGGSSLKTLEPELGTLQARLDRLLRRL QLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRAAHAILGGL<u>A</u>LTL<u>V</u>WAVRGLLL LKTRL

hIL-11 mutein deriving from 34aa-deleted native human hIL-11 -SEQ ID NO :12-:

PRAELDSTVLLTRSLLADTRQLAAQLRDKFPADGDHNLDSLPTLAMSAGALGA LQLPGVLTRLRADLLSYLRHVQWLRRAGGSSLKTLEPELGTLQARLDRLLRRL QLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRAAHAILGGL<u>Y</u>LTL<u>Y</u>WAVRGLLL LKTRL

hIL-11 mutein deriving from 34aa-deleted native human hIL-11 -SEQ ID NO :13-:

PRAELDSTVLLTRSLLADTRQLAAQLRDKFPADGDHNLDSLPTLAMSAGALGA LQLPGVLTRLRADLLSYLRHVQWLRRAGGSSLKTLEPELGTLQARLDRLLRRL QLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRAAHAILGGL<u>A</u>LTL<u>A</u>WAVRGLLL LKTRL

hIL-11 mutein deriving from 21aa-deleted native human hIL-11 -SEQ ID NO :14-:

PGPPPGPPRVSPDPRAELDSTVLLTRSLLADTRQLAAQLRDKFPADGDHNLDSL PTLAMSAGALGALQLPGVLTRLRADLLSYLRHVQWLRRAGGSSLKTLEPELGT LQARLDRLLRRLQLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRAAHAILGGL $\underline{\mathbf{X}}_1$ LTL $\underline{\mathbf{X}}_2$ WAVRGLLLLKTRL

wherein X1 and X2 are chosen from the group comprising:

- Alanine (A),
- Valine (V),
- Leucine (L),
- Isoleucine (I),
- Phenylalanine (F),
- Methionine (M),
- Proline (P),
- Tryptophan (W).

hIL-11 mutein deriving from 21aa-deleted native human hIL-11 -SEQ 1D NO :15-:

PGPPPGPPRVSPDPRAELDSTVLLTRSLLADTRQLAAQLRDKFPADGDHNLDSL PTLAMSAGALGALQLPGVLTRLRADLLSYLRHVQWLRRAGGSSLKTLEPELGT LQARLDRLLRRLQLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRAAHAILGGL<u>V</u>L TL<u>A</u>WAVRGLLLLKTRL

hIL-11 mutein deriving from 21aa-deleted native human hIL-11 -SEQ ID NO :16-:

PGPPPGPPRVSPDPRAELDSTVLLTRSLLADTRQLAAQLRDKFPADGDHNLDSL PTLAMSAGALGALQLPGVLTRLRADLLSYLRHVQWLRRAGGSSLKTLEPELGT LQARLDRLLRRLQLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRAAHAILGGL<u>A</u>L TLYWAVRGLLLLKTRL

hlL-11 mutein deriving from 21aa-deleted native human hlL-11 -SEQ ID NO :17-:

PGPPPGPPRVSPDPRAELDSTVLLTRSLLADTRQLAAQLRDKFPADGDHNLDSL PTLAMSAGALGALQLPGVLTRLRADLLSYLRHVQWLRRAGGSSLKTLEPELGT LQARLDRLLRRLQLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRAAHAILGGL<u>V</u>L TL<u>V</u>WAVRGLLLLKTRL

hIL-11 mutein deriving from 21aa-deleted native human hIL-11 -SEQ ID NO :18-:

PGPPPGPPRVSPDPRAELDSTVLLTRSLLADTRQLAAQLRDKFPADGDHNLDSL PTLAMSAGALGALQLPGVLTRLRADLLSYLRHVQWLRRAGGSSLKTLEPELGT LQARLDRLLRRLQLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRAAHAILGGL<u>A</u>L TL<u>A</u>WAVRGLLLLKTRL

hIL-11 mutein deriving from complete native human hIL-11 -SEQ ID NO :19-:

MNCVCRLVLVVLSLWPDTAVAPGPPPGPPRVSPDPRAELDSTVLLTRSLLADTR QLAAQLRDKFPADGDHNLDSLPTLAMSAGALGALQLPGVLTRLRADLLSYLRH VQWLRRAGGSSLKTLEPELGTLQARLDRLLRRLQLLMSRLALPQPPPDPPAPPL APPSSAWGGIRAAHAILGGLX,LTLX,WAVRGLLLLKTRL

wherein X1 and X2 are chosen from the group comprising:

- Alanine (A),
- Valine (V),
- Leucine (L),
- Isoleucine (I),
- Phenylalanine (F),
- Methionine (M),
- Proline (P),
- Tryptophan (W).

hlL-11 mutein deriving from complete native human hlL-11 -SEO ID NO :20-:

MNCVCRLVLVVLSLWPDTAVAPGPPPGPPRVSPDPRAELDSTVLLTRSLLADTR QLAAQLRDKFPADGDHNLDSLPTLAMSAGALGALQLPGVLTRLRADLLSYLRH VQWLRRAGGSSLKTLEPELGTLQARLDRLLRRLQLLMSRLALPQPPPDPPAPPL APPSSAWGGIRAAHAILGGL<u>V</u>LTL<u>A</u>WAVRGLLLLKTRL

hIL-11 mutein deriving from complete native human hIL-11 -SEO ID NO :21-:

MNCVCRLVLVVLSLWPDTAVAPGPPPGPPRVSPDPRAELDSTVLLTRSLLADTR QLAAQLRDKFPADGDHNLDSLPTLAMSAGALGALQLPGVLTRLRADLLSYLRH VQWLRRAGGSSLKTLEPELGTLQARLDRLLRRLQLLMSRLALPQPPPDPPAPPL APPSSAWGGIRAAHAILGGLALTLYWAVRGLLLLKTRL

hIL-11 mutein deriving from complete native human hIL-11 -SEQ ID NO :22-:

MNCVCRLVLVVLSLWPDTAVAPGPPPGPPRVSPDPRAELDSTVLLTRSLLADTR QLAAQLRDKFPADGDHNLDSLPTLAMSAGALGALQLPGVLTRLRADLLSYLRH VQWLRRAGGSSLKTLEPELGTLQARLDRLLRRLQLLMSRLALPQPPPDPPAPPL APPSSAWGGIRAAHAILGGL<u>V</u>LTL<u>V</u>WAVRGLLLLKTRL

hlL-11 mutein deriving from complete native human hIL-11 -SEQ ID NO :23-:

MNCVCRLVLVVLSLWPDTAVAPGPPPGPPRVSPDPRAELDSTVLLTRSLLADTR QLAAQLRDKFPADGDHNLDSLPTLAMSAGALGALQLPGVLTRLRADLLSYLRH VQWLRRAGGSSLKTLEPELGTLQARLDRLLRRLQLLMSRLALPQPPPDPPAPPL APPSSAWGGIRAAHAILGGL<u>A</u>UTL<u>A</u>WAVRGLLLLKTRL

IL-11 mutein deriving from 34aa-deleted native macaque IL-11 -SEQ ID NO:24-:

PRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ TRLDRLLRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG IRAAHAILGG L \underline{X}_1 LTL \underline{X}_2 WAVR GLLLLKTRL wherein X_1 and X_2 are chosen from the group comprising :

- Alanine (A),
- Valine (V),
- Leucine (L),
- Isoleucine (1),
- Phenylalanine (F),
- Methionine (M),
- Proline (P),
- Tryptophan (W).

IL-11 mutein deriving from 342a-deleted native macaque IL-11 -SEQ ID NO:25:

PRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH NLDSLPTLAM
SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ
TRLDRLLRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG
IRAAHAILGG LYLTLAWAVR GLLLLKTRL

IL-11 mutein deriving from 34aa-deleted native macaque IL-11 -SEQ ID NO:26:

PRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH NLDSLPTLAM
SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ
TRLDRLLRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG
IRAAHAILGG LALTLYWAVR GLLLLKTRL

IL-11 mutein deriving from 34aa-deleted native macaque IL-11 -SEQ ID NO:27:

PRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH NLDSLPTLAM
SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ
TRLDRLLRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG
IRAAHAILGG LVLTLWAVR GLLLLKTRL

IL-11 mutein deriving from 34aa-deleted native macaque IL-11 -SEQ ID NO:28-:
PRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH NLDSLPTLAM
SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ
TRLDRLLRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG
IRAAHAILGG LALTLAWAVR GLLLLKTRL

IL-11 mutein deriving from 21aa-deleted native macaque IL-11 -SEQ ID NO:29-:

PGPPPGSPR ASPDPRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ TRLDRLLRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG IRAAHAILGG L $\underline{\mathbf{X}}_1$ LTL $\underline{\mathbf{X}}_1$ WAVR GLLLLKTRL wherein X_1 and X_2 are chosen from the group comprising :

- Alanine (A),
- Valine (V),
- Leucine (L),
- Isoleucine (I),
- Phenylalanine (F),
- Methionine (M),
- Proline (P),
- Tryptophan (W).

IL-11 mutein deriving from 21aa-deleted native macaque IL-11 -SEQ ID NO:30-: PGPPPGSPR ASPDPRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ TRLDRLLRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG IRAAHAILGG LYLTLAWAVR GLLLLKTRL

IL-11 mutein deriving from 21aa-deleted native macaque IL-11 -SEQ ID NO:31-:
PGPPPGSPR ASPDPRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH
NLDSLPTLAM SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK
TLEPELGTLQ TRLDRLLRRL QLLMSRLALP QLPPDPPAPP
LAPPSSTWGG IRAAHAILGG LALTLYWAVR GLLLLKTRL

IL-11 mutein deriving from 2122-deleted native macaque IL-11 -SEQ ID NO:32-: PGPPPGSPR ASPDPRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ TRLDRLLRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG IRAAHAILGG LYLTLYWAVR GLLLLKTRL

IL-11 mutein deriving from 21aa-deleted native macaque IL-11 -SEQ ID NO:33-: PGPPPGSPR ASPDPRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ TRLDRLLRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG IRAAHAILGG LALTLAWAVR GLLLLKTRL

IL-11 mutein deriving from complete native macaque IL-11 -SEQ ID NO:34-:

MNCVCRLVLV VLSLWPDTAV APGPPPGSPR ASPDPRAELD STVLLTRSLL EDTRQLTIQL KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ TRLDRLLRRL QLLMSRLALP QLPPDPPAPP LAPPSSTWGG IRAAHAILGG LXLTLXLWAVR GLLLLKTRL

wherein X1 and X2 are chosen from the group comprising:

- Alanine (A),
- Valine (V),
- Leucine (L),
- Isoleucine (I),
- Phenylalanine (F),
- Methionine (M),
- Proline (P),
- Tryptophan (W).

IL-11 mutein deriving from complete native macaque IL-11 -SEQ ID NO:35-:

MNCVCRLVLV VLSLWPDTAV APGPPPGSPR ASPDPRAELD STVLLTRSLL
EDTRQLTIQL KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD
LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ TRLDRLLRRL QLLMSRLALP
QLPPDPPAPP LAPPSSTWGG IRAAHAILGG LYLTLAWAVR GLLLLKTRL

IL-11 mutein deriving from complete native macaque IL-11-SEQ ID NO:36-:
MNCVCRLVLV VLSLWPDTAV APGPPPGSPR ASPDPRAELD STVLLTRSLL
EDTRQLTIQL KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD
LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ TRLDRLLRRL QLLMSRLALP
QLPPDPPAPP LAPPSSTWGG IRAAHAILGG LALTLWAVR GLLLLKTRL

IL-11 mutein deriving from complete native macaque IL-11 -SEQ ID NO:37-:
MNCVCRLVLV VLSLWPDTAV APGPPPGSPR ASPDPRAELD STVLLTRSLL
EDTRQLTIQL KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD
LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ TRLDRLLRRL QLLMSRLALP
QLPPDPPAPP LAPPSSTWGG IRAAHAILGG LYLTLYWAVR GLLLLKTRL

IL-11 mutein deriving from complete native macaque IL-11-SEQ ID NO:38-:
MNCVCRLVLV VLSLWPDTAV APGPPPGSPR ASPDPRAELD STVLLTRSLL
EDTRQLTIQL KDKFPADGDH NLDSLPTLAM SAGALGALQL PSVLTRLRAD
LLSYLRHVQW LRRAMGSSLK TLEPELGTLQ TRLDRLLRRL QLLMSRLALP
QLPPDPPAPP LAPPSSTWGG IRAAHAILGG LALTLAWAVR GLLLLKTRL

IL-11 mutein deriving from 34aa-deleted native mouse IL-11 -SEQ ID NO:39-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG LX₁LTL_Xwavr GLLLLKTRL

wherein X1 and X2 are chosen from the group comprising:

- Alanine (A),
- Valine (V).
- Leucine (L),
- Isoleucine (I),
- Phenylalanine (F),
- Methionine (M),
- Proline (P),
- Tryptophan (W).

IL-11 mutein deriving from 34aa-deleted native mouse IL-11 -SEO ID NO:40-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG L<u>V</u>LTL<u>A</u>WAVR GLLLLKTRL

IL-11 mutein deriving from 34aa-deleted native mouse IL-11 -SEQ ID NO:41-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG LALTLWAVR GLLLLKTRL

IL-11 mutein deriving from 34aa-deleted native mouse IL-11 -SEQ ID NO:42-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG L<u>V</u>LTL<u>V</u>WAVR GLLLLKTRL

IL-11 mutein deriving from 34aa-deleted native mouse IL-11 -SEQ ID NO:43-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG LALTLAWAVR GLLLLKTRL

IL-11 mutein deriving from 21aa-deleted native mouse IL-11 -SEQ ID NO:44-:

PGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG LX1LTLX2WAVR GLLLLKTRL

wherein X1 and X2 are chosen from the group comprising:

- Alanine (A),
- Valine (V),
- Leucine (L),
- Isoleucine (I),
- Phenylalanine (F),
- Methionine (M),
- Proline (P),
- Tryptophan (W).

IL-11 mutein deriving from 21aa-deleted native mouse IL-11 -SEQ ID NO:45-:

PGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG L<u>V</u>LTL<u>A</u>WAVR GLLLLKTRL

IL-11 mutein deriving from 21aa-deleted native mouse IL-11 -SEQ ID NO:46-:

PGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG LALTLYWAVR GLLLLKTRL

IL-11 mutein deriving from 21aa-deleted native mouse IL-11-SEQ ID NO:47-:

PGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG L<u>V</u>LTL<u>V</u>WAVR GLLLLKTRL

IL-11 mutein deriving from 21aa-deleted native mouse IL-11 -SEQ ID NO:48-:

PGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG LALTLAWAVR GLLLLKTRL

IL-11 mutein deriving from complete native mouse IL-11 -SEQ ID NO:49-:

MNCVCRLVLV VLSLWPDRVV APGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG LX1LTLX2WAVR GLLLLKTRL

wherein X1 and X2 are chosen from the group comprising:

- Alanine (A),
- Valine (V),
- Leucine (L),
- Isoleucine (I),
- Phenylalanine (F),
- Methionine (M),
- Proline (P),
- Tryptophan (W).

IL-11 mutein deriving from complete native mouse IL-11 -SEQ ID NO:50-:

MNCVCRLVLV VLSLWPDRVV APGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG L<u>V</u>LTL<u>A</u>WAVR GLLLLKTRL

IL-11 mutein deriving from complete native mouse IL-11 -SEQ ID NO:51-:

MNCVCRLVLV VLSLWPDRVV APGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG LALTLVWAVR GLLLLKTRL

1L-11 mutein deriving from complete native mouse IL-11 -SEQ ID NO:52-:

MNCVCRLVLV VLSLWPDRVV APGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG L<u>V</u>LTL<u>V</u>WAVR GLLLLKTRL

IL-11 mutein deriving from complete native mouse IL-11 -SEQ ID NO:53-:

MNCVCRLVLV VLSLWPDRVV APGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH SLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYLRHVQW LRRAGGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPVIP LGPPASAWGS IRAAHAILGG LALTLAWAVR GLLLLKTRL

1L-11 mutein deriving from 34aa-deleted native rat IL-11 -SEO ID NO:54-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG LX,LTLX,WAVR GLLLLKTRL

wherein X1 and X2 are chosen from the group comprising:

- Alanine (A),
- Valine (V),
- Leucine (L),
- Isoleucine (I),
- Phenylalanine (F),
- Methionine (M),
- Proline (P),
- Tryptophan (W).

IL-11 mutein deriving from 34aa-deleted native rat IL-11 -SEO ID NO:55-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG L<u>V</u>LTL<u>A</u>WAVR GLLLLKTRL

IL-11 mutein deriving from 34aa-deleted native rat IL-11 -SEQ ID NO:56-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHA)LGG LALTLYWAVR GLLLLKTRL

IL-11 mutein deriving from 34aa-deleted native rat IL-11 -SEQ ID NO:57-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG L<u>V</u>LTL<u>V</u>WAVR GLLLLKTRL

IL-11 mutein deriving from 34aa-deleted native rat IL-11 -SEQ ID NO:58-:

PRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD ŁMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG LALTLAWAVR GLLLLKTRL

IL-11 mutein deriving from 21aa-deleted native rat IL-11 -SEQ ID NO:59-:

PGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG LX,LTLX,WAVR GLLLLKTRL

wherein X₁ and X₂ are chosen from the group comprising:

- Alanine (A),
- Valine (V),
- Leucine (L),
- Isoleucine (I),
- Phenylalanine (F).
- Methionine (M),
- Proline (P),
- Tryptophan (W).

IL-11 mutein deriving from 21aa-deleted native rat IL-11 -SEQ ID NO:60-:

PGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG L<u>y</u>LTL<u>A</u>WAVR GLLLLKTRL

IL-11 mutein deriving from 21aa-deleted native rat IL-11 -SEQ ID NO:61-:

PGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG L<u>A</u>LTL<u>Y</u>WAVR GLLLLKTRL

IL-11 mutein deriving from 21aa-deleted native rat IL-11 -SEQ 1D NO:62-:

PGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG L<u>Y</u>LTL<u>Y</u>WAVR GLLLLKTRL

IL-11 mutein deriving from 21aa-deleted native rat IL-11 -SEQ ID NO:63-:

PGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG L<u>A</u>LTL<u>A</u>WAVR GLLLLKTRL

IL-11 mutein deriving from complete native rat IL-11 -SEQ ID NO:64-:

MNCVCRLVLV VLSLWPDRVV APGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG L $\underline{\mathbf{X}}_1$ LTL $\underline{\mathbf{X}}_2$ WAVR GLLLLKTRL wherein X_1 and X_2 are chosen from the group comprising :

- Alanine (A),
- Valine (V),
- Leucine (L),
- Isoleucine (I),
- Phenylalanine (F),
- Methionine (M),
- Proline (P),
- Tryptophan (W).

IL-11 mutein deriving from complete native rat IL-11 -SEO ID NO:65-:

MNCVCRLVLV VLSLWPDRVV APGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG L<u>V</u>LTL<u>A</u>WAVR GLLLLKTRL

IL-11 mutein deriving from complete native rat IL-11 -SEQ ID NO:66-:

MNCVCRLVLV VLSLWPDRVV APGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG LALTLYWAVR GLLLLKTRL

IL-11 mutein deriving from complete native rat IL-11 -SEQ ID NO:67-:

MNCVCRLVLV VLSLWPDRVV APGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG L<u>V</u>LTL<u>V</u>WAVR GLLLLKTRL

IL-11 mutein deriving from complete native rat IL-11 -SEQ ID NO:68-:

MNCVCRLVLV VLSLWPDRVV APGPPAGSPR VSSDPRADLD SAVLLTRSLL ADTRQLAAQM RDKFPADGDH NLDSLPTLAM SAGTLGSLQL PGVLTRLRVD LMSYFRHVQW LRRAAGPSLK TLEPELGALQ ARLERLLRRL QLLMSRLALP QAAPDQPAVP LGPPASAWGS IRAAHAILGG L<u>A</u>LTL<u>A</u>WAVR GLLLLKTRL

Joined CDS for human complete native IL-11 -SEQ ID NO:69-:

Joined CDS for the IL-11 mutein which derives from the 34aa-deleted human IL-11 - SEQ ID NO:70-:

wherein the codon $n_1n_2n_3$ and the codon $n_4n_5n_6$ are both chosen among the group comprising the nucleotide codons which codes for a hydrophobic aminoacid, namely for Alanine (A), Valine (V), Leucine (L), Isoleucine (I), Phenylalanine (F), Methionine (M), Proline (P), Tryptophan (W).

 $n_1n_2n_3$ and $n_4n_5n_6$ can be chosen among the group comprising the following nucleotide codons:

- GCT, GCC, GCA, GCG
- GTT, GTC, GTA, GTG,
- TTA, TTG, CTT, CTC, CTA, CTG,
- ATT, ATC, ATA,
- TTT, TTC,
- ATG,
- CCT, CCC, CCA, CCG,
- TGG.

FIGURE 16A

Joined CDS for the IL-11 mutein which derives from the 21aa-deleted human IL-11 - SEQ ID NO:71-:

wherein the codon $n_1n_2n_3$ and the codon $n_4n_5n_6$ are as defined in Figure 16A.

Joined CDS for the IL-11 mutein which derives from the complete human IL-11 -SEQ ID NO:72-:

wherein the codon n₁n₂n₃ and the codon n₄n₅n₆ are as defined in Figure 16A.

FIGURE 16B

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Mutated AY207429 nucleic acid -SEQ ID NO:74-:

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1 acacctgtat teceaceact ttgggagget gaggegggag gatgaeetga geteaggagt
  61 ttgagaccag cctgggcaac atggcaaaac cctatctcta ctaaaaatac aaaaaatagc
 121 caggcatggt ggcgggtgcc tgtaatccca gctactcagg aggctgaggc atgagaatca
 181 cttgaacctg ggaggcggag gttacagtga gctgagatca caccactgca ccccagcctg
 241 ggtgacacag cgagactotg totoaaaaaa accaaaaacg aggccaggca cggtagotoa
 301 cacctgtcat cccagcactt tgggaggccg aggcaggcgg atcacgaagt caggagttcg
 361 agaccagect ggccaacatg gtaagacccc gtetetacta aaaatacaaa attagccggg
 421 tgtggtggcg cacacctgta atcccagcta cttgggaggc tgaggcagga gaatcgcttg
 481 aaccogggag gtggaggttg cagtgagctg agattgtgcc attgatcgcg ccattgcact
 541 ccagcctggg tgacagagtg agactcagta ccaaaaaaca aacaaacaaa aaacaaacaa
 601 aaaatgagaa aggettttae tetetgeece cattgetgag teeceaacat eteagegtet
 661 etgtetttet aatatetetg teteceettt tetgteeetg gggeetetee gteeetgtea
 721 etetgecceg tgtetetgtt tgeetggtge etttetteag etgeggeate etetgtetea
 781 gagtettggt gtetetgtte ettteecete ggggtetece tgggtetece caagteecte
 841 etgetgtett cetecegete tetgatetet gacteceaga acetetecet etgtetecag
901 ggetgeecet etgatectet ttgettetet ggtgtgtete tetggetgee tecatetetg
 961 tggatctccg tctccctgtc tctgtctcag tctgtccttc actctgtgtg tgtgtgtgt
1021 tgtctctctc tctctctctc cttcccttcc actccctctt cctcctgcct ccacctctcc
1081 aggcccetgt cttgtccctc cgtccggcct ttctctgcct ttccgtcctc ctgcctcccc
1141 atotototot gotagtoctg gtocagoogg accoccacco acagtogggo cocagogott
1201 gagcctgagt gtctgctccg gcccgtggag gtggagggag gggacgccaa tgacctcacc
1261 ageccetete egaceacece eccettece titteaactt ticeaacttt teetteegtg
1321 coetectorg agegeggegg egtgageeet geaaggeage egeteegtet gaatggaaaa
1381 ggcaggcagg gagggtgagt caggatgtgt caggccgccc teccetgccg cetgcccccc
1441 gcccgcccgc cccagccccc tatataaccc cccaggcgtc cacactccct cactgccgcg
1501 gecetgetge teagggeaca tgeeteeeet eeceaggeeg eggeecaget gacceteggg
1561 geteccegg cageggacag ggaagggtta aaggeeeeeg getecetgee ecetgeeetg
1621 gggaacccct ggccctgtgg ggacatgaac tgtaagttgg ttcatgggga gggtggaggg
1681 gacagggagg cagggaggag agggacccac ggcgggggtg ggagcagacc ccgctgagtc
1741 gcacagagag ggacccggag acaggcagcc ggggaggaga gcagcttcgg agacaggagg
1801 cggcggagga gatgggcaga gagagacaca gacaggagcg gatggaggca gccaatcaga
1861 ggcgccgcag gagggacggg ccagacaggg ccccgagagg gagcgagacg cggagaccga
1921 gcaggggcag ggacgcaggg actggtgccg ggagggaggt gacccccatc gacccaggcc
1981 ccagggagcc cgcggggacc gggagactcc ctgggattcc ggcagagagg ctccggaggg
2041 aaactgagge agggteegeg gagageggag caagecaggg agtagegace ecageegggg
2101 ggaggagaga gactgggcgc ggggggaaag cggggagagc cgggcagatg cggccgacgg
2161 aggcgcggac agaccgacgg ctggcgggcc cggggggcgg gctgggggtg tgcgaggcgc
2221 gggcggccgg ggagcgctga ttggctggcg ggtggccggg tgggcggggc ggccggggtg
2281 ggctgcgggg agcgagctcc ggacccccgc gccccccgcg cccccgcgc cccccgccgc
2341 cagetetece getecegegg eccaggeggg cecatggete tgeceetete egeceaggtg
2401 cgctgcggcc cgggcttctg ccgcccaccc ggcggggctc ctgggagggc gtctaagggg
2461 tetecegtgg gagaggteeg tgteteeegg geteegteet ggettetgge teetteeeet
2521 gctcccagec agctcgggct cccgcggccc ggggaggggg caggttctgg cctgtgcctc
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2641 eggecegtet accegeceeg ggecgegtet getecgaegg geggggeage eagageeagg
2701 gagggagagg gaagcccgcc tggccctgcg acctgcccgc gggcgttcca ccctgggact
2761 taagacetee agetecatee teectaagge egggagteea ggeeceagae eeteeteece
2821 gagacccagg agtecagacc ccaggeette eteceteaga cetaggagte caggeeccca
2881 geeteteete eeteagaeee aggaggagte cagaeeeeag tteeteetee eteagaeeeg
2941 ggagtccagg cccaggccct cctctctcag acccggagtc cagcctgagc tctctgcctt
3001 accetgeece caggigiting cogeetygic etygicging transcripting gecagataca
3061 getgtegeee etgggeeaee acetggeece cetegagttt ecceagacee tegggeegag
3121 ctggacagea ccgtgctcct gacccgctct ctcctggcgg acacgcggca gctggctgca
3181 cagctggtag gagagactgg gctggggcca gcacaggagt gagaggcaga gaggaacgga
```

```
3241 gaggagtetg egggeageea ettggagggg ttetgggete teaggtggea gagtgaggga
3301 ggggaagagt tgggggcctg gcgtggggga tggagggagc cccgaggctg ggcaggggcc
3361 acctcacage ttttttccct gccagaggga caaattccca gctgacgggg accacaacct
3421 ggattccctg cccaccctgg ccatgagtgc gggggcactg ggagctctac aggtaagggc
3481 aagggagtgg gctggggaca aggtgggagg caggcagtga aggggggggg gaggatgagg
3541 ggcactggtc gggtgttctc tgatgtcccg gctctatccc cagctcccag gtgtgctgac
3601 aaggetgega geggaeetae tgteetaeet geggeaegtg eagtggetge geegggeagg
3661 tggctcttcc ctgaagaccc tggagcccga gctgggcacc ctgcaggccc gactggaccg
3721 getgetgege eggetgeage teetggtatg teetggeece aagacetgae acceeagace
3781 occaeccetg geoceaaaat cetgtggeet gagteettga ageetgagae ceeagaeceg
3841 agtgcaacag ccccgctctg agaccctgac accctaacag cccgctctga gaccctgaca
3901 ccgtaacage cccgetetga gaccetgace etaacagtee tgetetgaga ccetgaceet
3961 gcagtcccaa gatcctgtgg ccctgagacc ctgaggccct agacccccaa atcctgccca
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4081 agateccage ecetaagace caagacecca teetgaagee caaageettg agaatteaaa
4141 tecteacete aagaettgga gaeeetggee eeatgaeatt gaaaaceatg gaeetggeea
4201 ggcgtggtgg ctcacgcctg taatcccagc actttgggag gccgaggcaa gtggatcacc
4261 tgaggtcggg agttcaagac cagccagacc aacatggtga aaccctgtct ctactaaaaa
4321 tacaaaatta gccaggcgtg gtggtgcatg cctgtaatcc cagctacttg ggaggctgag
4381 gcaggagaat cgcttgaacc tgggaggcgg aggttgcagt gagccgagat cgcaccatta
4441 cactocagee tgggeaacaa gageaaaact ecetetete caaaaaaaaa aaaaaaaaaa
4501 aaaagaagga aaagaaaacc atggacctcc agaccctgag accccaggcc ccagccctga
4561 gatectgaca tettaaagat eecaggeeet aagatacaag acettgacee aaageeagee
4621 ttgggaccct ggctgtacaa acccaagacc tccaggacct agaccccgag ccctgaggcc
4681 ctatgtetca eteceaacat egaaaaceet gacaceteag ateetgagee tgegeetgta
4741 cgactccaag acceteaett ccaaagecag geccaaagec etgagaccag aagaettcaa
4801 accotggtto ttgggcctaa ctccaaagac cotggatoto aaattocaac ttotagotot
4861 gagactccag ccctcaccca tgagttcctg aacttgaacc cagagacccc atctctaaga
4921 cttcagcctt gagatccagg gcctgaccct agactcgagc ccacagacct cagatactgt
4981 ctgtaaaacc ccagctctgg tggggagcag tggctcactc ctgtaatccc aaggcagggg
5041 aggccaaggc agaaggacct cttgaggcca tgagtttgag acagcctggg cagcatagca
5101 agactotgtt tottaattat tattattatt attattttt ggagacagag totogogoto
5161 tgttgcccag gctagagtgc aatggtgcca tttcggcttg ctggaacctc cgcctcetgg
5221 gctcaagcga ttetcetgee teagestest gagtagetgg gaetteaggt gcacactges
5281 acacceggat aattititig tattitagta gacacagggt ticaccgtgt tgeccagget
5341 ggtcacaaac tcctgagctc aggccatccg cccgcctcgg cctcccaaag cgctgggata
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5581 tggaageceg teetgactee agenteeatt tteggaacee cacagectga agageteeeg
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5761 ctetectete eccaeagatg tecegeetgg ecctgeecea gecaeeceeg gaeeegeegg
5821 egecceget ggegececc tecteageet gggggggcat cagggegge caegecatec
5881 tgggggggct gmmmctgaca cttmmmtggg ccgtgagggg actgctgctg ctgaagacte
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6001 tttatttcag tactgggggc gaaacagcca ggtgatcccc ccgccattat ctcccctag
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6121 ttcaggagca ggggtgggag gcaggtggac tcctgggtcc ccgaggagga ggggactggg
6181 gtcccggatt cttgggtctc caagaagtct gtccacagac ttctgccctg gctcttcccc
6241 atctaggcct gggcaggaac atatattatt tatttaagca attacttttc atgttggggt
6301 ggggacggag gggaaaggga agcctgggtt tttgtacaaa aatgtgagaa acctttgtga
6361 gacagagaac agggaattaa atgtgtcata catatccact tgagggcgat ttgtctgaga
6421 gctggggctg gatgcttggg taactggggc agggcaggtg gaggggagac ctccattcag
6481 gtggaggtcc cgagtgggcg gggcagcgac tgggagatgg gtcggtcacc cagacagctc
6541 tgtggaggca gggtctgagc cttgcctggg gccccgcact gcatagggcc gtttgtttgt
                           FIGURE 17
```

```
6601 tttttgagat ggagtctcgc tctgttgcct aggctggagt gcagtgaggc aatctaaggt
6661 cactgcaacc tccacctccc gggttcaagc aattctcctg cctcagcctc ccgattagct
6721 gggatcacag gtgtgcacca ccatgcccag ctaattattt atttcttttg tatttttagt
6781 agagacaggg tttcaccatg ttggccaggc tggtttcgaa ctcctgacet caggtgatec
6841 tectgeeteg geeteecaaa gtgetgggat tacaggtgtg ageeaccaca cetgacccat
6901 aggtottoaa taaatattta atggaaggtt ccacaagtoa cootgtgato aacagtacco
6961 gtatgggaca aagetgcaag gtcaagatgg ttcattatgg ctgtgttcac catagcaaac
7021 tggaaacaat ctagatatcc aacagtgagg gttaagcaac atggtgcatc tgtggataga
7081 acgccaccca gccgcccgga gcagggactg tcattcaggg aggctaagga gagaggcttg
7141 cttgggatat agaaagatat cctgacattg gccaggcatg gtggctcacg cctgtaatcc
7201 tggcactttg ggaggacgaa gcgagtggat cactgaagtc caagagtttg agaccggcct
7261 gcgagacatg gcaaaaccct gtctcaaaaa agaaagaatg atgtcctgac atgaaacagc
7321 aggctacaaa accactgcat gctgtgatcc caattttgtg tttttctttc tatatatgga
7381 ttaaaacaaa aatcctaaag ggaaatacgc caaaatgttg acaatgactg tctccaggtc
7441 aaaggagaga ggtgggattg tgggtgactt ttaatgtgta tgattgtctg tattttacag
7501 aatttctgcc atgactgtgt attttgcatg acacatttta aaaataataa acactatttt
7561 tagaataaca gaatatcagc ctcctcctct ccaaaaataa gccctcagga ggggacaaag
7621 ttgaccgctg attgagcctg tcagggctgt gcactaagtg tgggcttttt acttacacaa
7681 tecteetgga etettgaata egecetgttt tacaggegag ggaaactgag teteagacaa
7741 ggagtgggga ctctgttgca caaagtcaca cagctaggga gaggtggaag tgggattctg
7801 cgccgtgtct ggctctttcc caaagctctc tttgcaagtc ggtgttgagg aatcctcgcc
7861 acatgcacac acatgagata tggagaaaca ggttcagtaa ggatttgggt cttacccagg
7921 gcctagagaa gggtcaatgg cagagtaggg atgataattc aaatgcttta gttacttttc
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8041 ggttggaggt gcccatcetg ggcccggagt tttgattcac ccatcatagc cctcaagact
8101 ccaggetgge tgggegeggt ggeteaegee tgtaateeea geaetttggg aggetgagge
8161 gggtggatca cttgaggtca ggagttcaag gccagcctga ccaacatgga gaaaccctgt
8221 ctctactaaa aatacaatcc agctactcgg aaggctgagg caggagaatc gctcgaaccc
8281 aggagacggg ggttgcggtg agccgagatc acatcacaaa cagccctagg cagtgcgggg
8341 ccccaggega ggctcagacc tgcctccaca gagctgtctg ggtgatcgtg cctcctccgt
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8461 atggagtete getetgttge etaggetgga gtgeagtgtg geaatetaag eteactgeet
8521 gggcaacaag agtgaaattc catctcaaaa aacaaaaaac aaacaaacaa acaaaaaact
8581 ccaggctgta tccctggagg agaagggagc ccacagtccc cggagagttc ctggaagagg
8641 cccctgtgtg tccgatgagg tcacaaagcc cctccaccag aggctcctcc cccagacccc
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8761 acgctctgta acgctgagct ccaggcaccc gtgaagcccc acgggtcaag gctggtgggc
8821 cggggctggg aggcctgcac gcctgggttc tgggtcccta aaccagtacc catccaccac
8881 agccaccatg atctggcttc gaaacaggag gtgccttgag ccgctccagg gcaccccgaa
8941 gtgggtccct gttctggggg agctgcaaaa gaccctccag aagggcgagt acctgcccct
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9061 ggctgagagg gacagagggg aagcaaggcc ccccgtgctg ggggatcttg agagggaacg
9121 ggatttagca gtcactgtgt gggggacgat caggagggag gctcaggctg tggctgctgg
9181 aggaaggagt ggtcccagcc ccctctccct ggctgcccca ggtgacccat caagggggcc
9241 cagtgttcgt gaatcacaga accaaccggc tggccatggg cgtggccgcc tccctgccag
9301 gcctggtgtt gcctgacatc ttgctgatcg gccagcccgc cgaggacagg gactgctccg
9361 gcctcgtgct gaccaggtgc cgcatccccc aacccctcgg ccgccccctc cacccctcct
9421 gctctagacg ctcccctctc cctctcccag gatgatcccc ctggacctcg tccacctctg
9481 cgtccatgac ctctctgcct ggcgcctgaa gctgcgcctg gtctcgggcc gccagtacta
9541 cctggccctg gacgcccctg acaacgaggt gggcttcctg ttccactgct gggtccgcct
9601 catcaacctg cttcaggage cggctcccac ctggaccccc aggaccacge gcacggcccc
9661 cctggatatg ccgctggcca aagcgcctgc ctccacctgg cacctgcagg tgggatccca
9721 getecacaga ecagggeatg geaggeecea ggaaccetee ggeeagatee agaggggaet
9781 cgaccaagag cccaaagtct agg
```

wherein the codon n₁n₂n₃ and the codon n₄n₅n₆ are as defined in Figure 16A.

mRNA of IL-11 mutein deriving from human IL-11 -SEO ID NO:75-:

gaa ggg uua aag gcc ccc ggc ucc cug ccc ccu gcc cug ggg aac ccc ugg ccc ugu ggg gac aug aac ugu guu ugc cgc cug guc cug guc gug cug agc cug ugg cca gau aca gcu guc gcc ccu ggg cea cea ceu gge cee ceu ega guu uce cea gae ceu egg gee gag eug gae age ace gug euc eug ace ege ueu eue eug geg gae aeg egg eag eug geu gea eag eug agg gae aaa uue eea geu gae ggg gac cac aac cug gau ucc cug ccc acc cug gcc aug agu gcg ggg gca cug gga gcu cua cag cuc cca ggu gug cug aca agg cug cga gcg gac cua cug ucc uac cug cgg cac gug cag ugg cug cgc egg gea ggu gge ueu uee eug aag ace eug gag eee gag eug gge ace eug eag gee ega eug gae cgg cug cug cgc cgg cug cag cuc cug aug ucc cgc cug gcc cug ccc cag cca ccc ccg gac ccg ccg geg eec eeg eug geg eec eec uec uez gee ugg ggg gge auc agg gee gee eac gee auc eug ggg ggg cug ninin cug aca cuu ninin ugg gcc gug agg gga cug cug cug cug aag acu cgg cug uga ccc ggg gcc caa age cac cae cgu ccu ucc aaa gcc aga ucu uau uua uuu auu uau uuc agu acu ggg ggc gaa aca gcc agg uga ucc ccc cgc cau uau cuc ccc cua guu aga gac agu ccu ucc gug agg ccu ggg ggg cau cug ugc cuu ann nan acu nan nan cag gag cag ggg ngg gag gca ggu gga cuc cug ggu ccc cga gga ggg gac ugg ggu ccc gga uuc uug ggu cuc caa gaa guc ugu cca cag acu ucu gcc cug gcu cuu ccc cau cua ggc cug ggc agg aac aua uau uau uua uuu aag caa uua cuu uuc aug uug ggg ugg gga cgg agg gga aag gga agc cug ggu uuu ugu aca aaa aug uga gaa acc uuu gug aga cag aga aca ggg aan naa aug ugu can aca nan cca cuu gag ggc gau und nen dad age neg dae end and enn des nas end des cas des age neg des dae ene can uca ggu gga ggu ccc gag ugg gcg ggg cag cga cug gga gau ggg ucg guc acc cag aca gcu cug ugg agg cag ggu cug agc cuu gcc ugg ggc ccc gca cug cau agg gcc guu ugu uuu uuu gag aug gag ucu cgc ucu guu gcc uag gcu gga gug cag uga ggc aau cua agg uca cug caa ccu cca ccu ccc ggg uuc aag caa uuc ucc ugc cuc agc cuc ccg auu agc ugg gau cac agg ugu gca cca cca ugc cca gcu aau uau uua uuu cuu uug uau uuu uag uag aga cag ggu uuc acc aug uug gcc agg cug guu ucg aac ucc uga ccu cag gug auc cuc cug ccu cgg ccu ccc aaa gug cug gga uua cag gug uga gcc acc aca ccu gac cca uag guc uuc aau aaa uau uua aug gaa ggu ucc aca agu cac ccu gug auc aac agu acc cgu aug gga caa gcu gca agg uca aga ugg uuc auu aug gcu gug uuc acc ana gca aac ugg aaa caa ucu aga uau cca aca gug agg guu aag caa can ggu gca ucu gug

gau aga acg cca ccc agc cgc ccg gag cag gga cug uca uuc agg gag gcu aag gag aga ggc uug cuu ggg aua uag aaa gau auc cug aca uug gcc agg cau ggu gge uca cgc cug uaa ucc ugg cac uuu ggg aga acg aag cga gug gau cac uga agu cca aga guu uga gac cgg ccu gcg aga cau ggc aaa acc cug ucu caa aaa aga aag aau gau guc cug aca uga aac agc agg cua caa aac cac ugc aug cug uga ucc caa auu guu ugu guu uuu cuu ucu aua uau gga uua aaa caa aaa ucc uaa agg gaa aua cgc caa aau guu gac aau gac ugu cuc cag guc aaa gga gag agg ugg gau ugu ggg uga cuu uua aug ugu ugu aug aug aca cau uuu aaa aau aau aaa cac uau uuu uag aau

wherein the codon $n_1n_2n_3$ and the codon $n_4n_5n_6$ are both chosen among the group comprising the nucleotide codons which codes for a hydrophobic aminoacid, namely for Alanine (A), Valine (V), Leucine (L), Isoleucine (I), Phenylalanine (F), Methionine (M), Proline (P), Tryptophan (W).

 $n_1n_2n_3$ and $n_4n_5n_6$ can be chosen among the group comprising the following nucleotide codons:

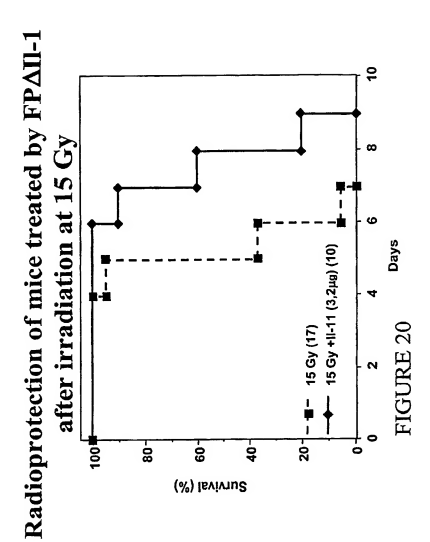
- GCU, GCC, GCA, GCG
- GUU, GUC, GUA, GUG,
- UUA, UUG, CUU, CUC, CUA, CUG,
- AUU, AUC, AUA,
- UUU, UUC,
- AUG,
- CCU, CCC, CCA, CCG,
- UGG.

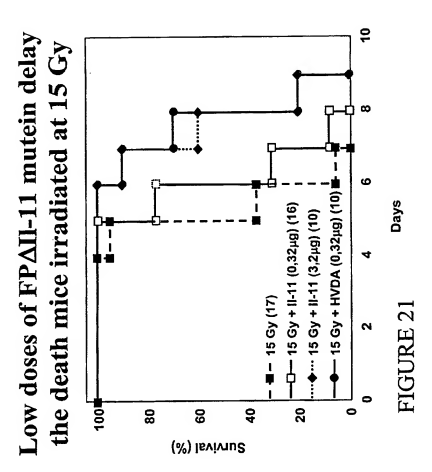
Gene of IL-11 muteins deriving from human IL-11 - SEQ ID NO:76-:

```
gggaacccct ggccctgtgg ggacatgaac tgtaagttgg ttcatgggga gggtggaggg
gacagggagg cagggaggag agggacccac ggcgggggtg ggagcagacc ccgctgagtc
gaacagagag ggacccggag acaggcagcc ggggaggaga gcagcttcgg agacaggagg
cggcggagga gatgggcaga gagagacaca gacaggagcg gatggaggca gccaatcaga
ggcgccgcag gagggacggg ccagacaggg ccccgagagg gagcgagacg cggagaccga
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ccagggagcc cgcggggacc gggagactcc ctgggattcc ggcagagagg ctccggaggg
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ggaggagaga gactgggcgc ggggggaaag cggggagagc cgggcagatg cggccgacgg
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accotggite tigggeetaa ciccaaagae ceiggatete aaattecaae tietagetet
gagactocag coctcaccca tgagttcctg aacttgaacc cagagacccc atctctaaga
cttcagcctt gagatccagg gcctgaccct agactcgagc ccacagacct cagatactgt
```

ctgtaaaacc ccagctctgg tggggagcag tggctcactc ctgtaatccc aaggcagggg aggccaaggc agaaggacct cttgaggcca tgagtttgag acagcctggg cagcatagca agactotgtt tottaattat tattattatt attattttt ggagacagag totogcgcto tgttgcccag gctagagtgc aatggtgcca tttcggcttg ctggaacctc cgcctcctgg getcaagega tteteetgee teageeteet gagtagetgg gaetteaggt geacactgee acacceggat aattittig tattitagta gacacagggt ticacegtgt tgcccagget ggtcacaaac tcctgagctc aggccatecg cccgcctcgg cctcccaaag cgctgggata acaggogtga tocogogogo otggottott aattgttota acagoagoca caacaacaaa aacccagete tgagatteca geeceggega etetaacagt eccaggeeeg ateceteaee tagaaccgag atgccagccc tgactccaca gacttcaccc ccaaccccca cactcagctc tggaageecg teetgaetee ageeteeatt tteggaaeee cacageetga agageteeeg gectaaacac tteaceceae gegecacagt ecceetgtga atatgeagee ecgatteage tgcageteca cagcaccect gccctgcacc cccgctgcac cccctacctg tgactcacct ctctcctctc cccacagatg tcccgcctgg ccctgcccca gccacccccg gacccgccgg egececeget ggegeeecce tecteageet gggggggeat cagggeegee caegecatee tgggggggct gniningctgaca cttniningtggg ccgtgagggg actgctgctg ctgaagactc ggctgtgacc cggggcccaa agccaccacc gtccttccaa agccagatct tatttattta tttatttcag tactgggggc gaaacagcca ggtgatcccc ccgccattat ctccccctag ttagagacag tccttccgtg aggcctgggg ggcatctgtg ccttatttat acttatttat ttcaggagca ggggtgggag gcaggtggac tcctgggtcc ccgaggagga ggggactggg gtcccggatt cttgggtctc caagaagtct gtccacagac ttctgccctg gctcttcccc atctaggect gggcaggaac atatattatt tatttaagca attactttte atgttggggt ggggacggag gggaaaggga agcctgggtt tttgtacaaa aatgtgagaa acctttgtga gacagagaac agggaattaa atgtgtcata catatccact tgagggcgat ttgtctgaga getggggetg gatgettggg taactgggge agggcaggtg gaggggagac etecatteag gtggaggtcc cgagtgggcg gggcagcgac tgggagatgg gtcggtcacc cagacagctc tgtggaggca gggtctgagc cttgcctggg gccccgcact gcatagggcc gtttgttgt tttttgagat ggagtctcgc tctgttgcct aggctggagt gcagtgaggc aatctaaggt cactgcaacc tccacctccc gggttcaagc aattotootg cotcagcotc ccgattagct gggatcacag gtgtgcacca ccatgcccag ctaattattt atttcttttg tatttttagt agagacaggg tttcaccatg ttggccaggc tggtttcgaa ctcctgacct caggtgatcc tectgeeteg geeteecaaa gtgetgggat tacaggtgtg agceaceaca cetgaeceat aggicticaa taaatattia atggaaggit ccacaagica ccctgtgatc aacagtaccc gtatgggaca aagctgcaag gtcaagatgg ttcattatgg ctgtgttcac catagcaaac tggaaacaat ctagatatcc aacagtgagg gttaagcaac atggtgcatc tgtggataga acgccaccca gccgcccgga gcagggactg tcattcaggg aggctaagga gagaggcttg cttgggatat agaaagatat cctgacattg gccaggcatg gtggctcacg cctgtaatcc tggcactttg ggaggacgaa gcgagtggat cactgaagtc caagagtttg agaccggcct gcgagacatg gcaaaaccct gtctcaaaaa agaaagaatg atgtcctgac atgaaacagc aggetacaaa accactgcat getgtgatce caattttgtg tttttettte tatatatgga ttaaaacaaa aatoctaaag ggaaatacgo caaaatgttg acaatgactg totocaggto aaaggagaga ggtgggattg tgggtgactt ttaatgtgta tgattgtctg tattttacag aatttctgcc atgactgtgt attttgcatg acacatttta aaaataataa acactatttt tagaat

wherein the codon n₁n₂n₃ and the codon n₄n₅n₆ are as defined in Figure 16A.





Parental (non-mutated) nucleotide sequence FPAIL-11 = SEQ ID NO:77 =

Parental (non-mutated) amino acid sequence of FP Δ IL-11 = SEQ ID NO:78 = MDYKDDDDKEGRRASVASPDPRAELDSTVLLTRSLLADTRQLAAQLRDKFPA DGDHNLDSLPTLAMSAGALGALQLPGVLTRLRADLLSYLRHVQWLRRAGGSS LKTLEPELGTLQARLDRLLRRLQLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRA AHAILGGL \underline{H} LTL \underline{D} WAVRGLLLLKTRL

Mutated amino acid sequence of FPAIL-11 = SEQ ID NO:80 of the invention = MDYKDDDDKEGRRASVASPDPRAELDSTVLLTRSLLADTRQLAAQLRDKFPADGDHNLDSLPTLAMSAGALGALQLPGVLTRLRADLLSYLRHVQWLRRAGGSSLKTLEPELGTLQARLDRLLRRLQLLMSRLALPQPPPDPPAPPLAPPSSAWGGIRAAHAILGGLYLTLAWAVRGLLLLKTRL

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Primers used for inverse PCR mutagenesis of FPAIL-11:

Muteins	Primers
H182/V	G422 pACACTTGACTGGGCCGTACGGGGAC (s) SEQ ID NO:81
	G412 pCAGAACCAGCCCCCCAGGATGG (as) SEQ ID NO:82
D186/V	G410 pACACTTGTCTGGGCCGTACGGGGAC (s) SEQ ID NO:83
	G421 pCAGGTGCAGCCCCCCAGGATGG (as) SEQ ID NO:84
D186/A	G411 pACACTTGCCTGGGCCGTACGGGGAC (s) SEQ ID NO:85
	G421 pCAGGTGCAGCCCCCCAGGATGG (as) SEQ ID NO:86
H182/V-	G410 pACACTTGTCTGGGCCGTACGGGGAC (s) SEQ ID NO:87
D186/V	G412 pCAGAACCAGCCCCCCAGGATGG (as) SEQ ID NO:88
H182/V-	G411 pACACTTGCCTGGGCCGTACGGGGAC (s) SEQ ID NO:89
D186/A	G412 pCAGAACCAGCCCCCCAGGATGG (as) SEQ ID NO:90

FIGURE 24

AB AB AB Site III Dies site I

Figure 25A

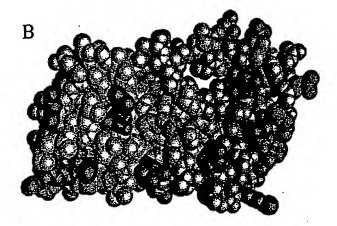


Figure 25B

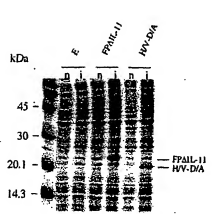


Figure 26

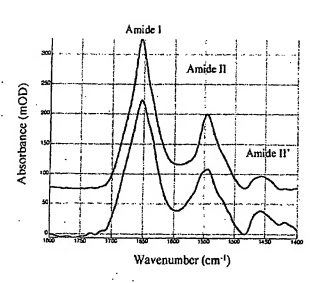


Figure 27

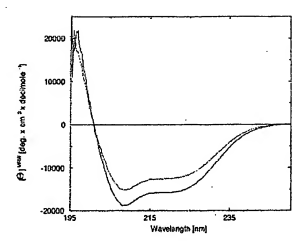


Figure 28

•

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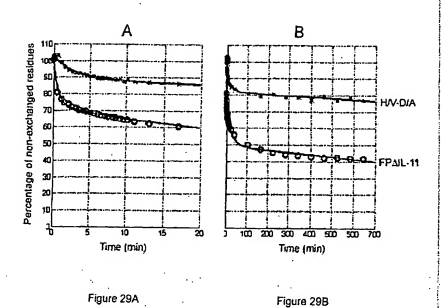
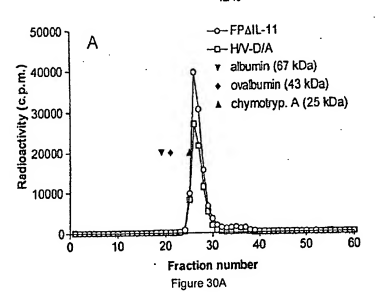


Figure 29B



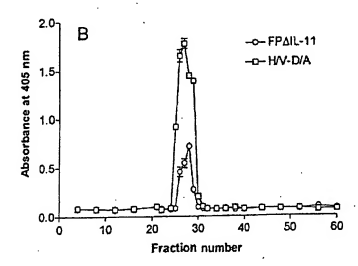
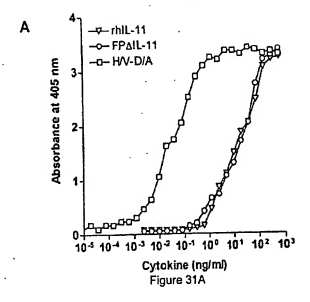


Figure 30B

the state of the s



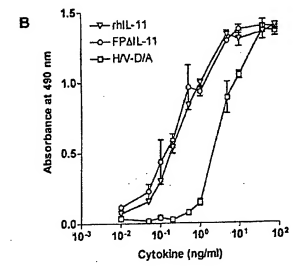


Figure 31B

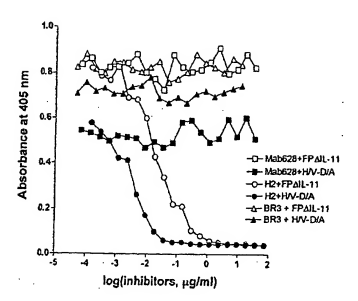


Figure 32

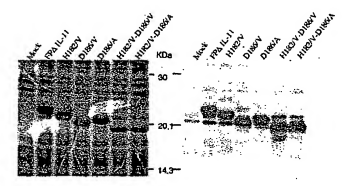


Figure 33

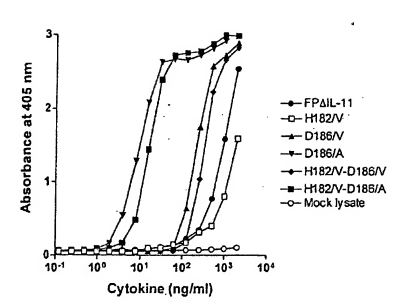


Figure 34

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PATENT COOPERATION TREATY

	From the INTERNATIONAL BUREAU		
PCT	То:		
NOTIFICATION OF THE RECORDING OF A CHANGE (PCT Rule 92bis.1 and Administrative Instructions, Section 422) Date of mailing (day/month/year) 08 July 2005 (08.07.2005)	PARIS, Fabienne Ernest Gutmann - Yves Plasseraud S.A 3 Rue Chauveau-Lagarde F-75008 Paris France		
Applicant's or agent's file reference FP/FFC-PARIS	IMPORTANT NOTIFICATION		
International application No. PCT/EP2004/009165	International filing date (day/month/year) 29 July 2004 (29.07.2004)		
1. The following indications appeared on record concerning: the applicant the inventor Name and Address PARIS, Fabienne Cabinet Plasseraud 65/67, rue de la Victoire F-75440 Paris Cedex 09 France	X the agent the common representative State of Nationality State of Residence Telephone No. +33 1 40 16 70 00 Facsimile No. +33 1 42 80 01 59 Teleprinter No.		
The International Bureau hereby notifies the applicant that the person the name X the additional than the person The name X the additional than the person The name X the additional than the person The name			
Name and Address PARIS, Fabienne Ernest Gutmann - Yves Plasseraud S.A 3 Rue Chauveau-Lagarde F-75008 Paris France 1 8. 07. 2	State of Nationality Telephone No. +33 1 44 51 18 00 Facsimile No. +33 1 42 66 08 90 Teleprinter No.		
3. Further observations, if necessary: TEAM 14			
4. A copy of this notification has been sent to: X the receiving Office the International Searching Authority the International Preliminary Examining Authority	X the designated Offices concerned the elected Offices concerned other:		
The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland Facsimile No. (41-22) 338.89.70	Isabella BULOZ (Fax 338-87.20) Telephone No. (41-22) 338 8479		

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